

World Health Organization Model List of Essential In Vitro Diagnostics

First edition (2018)

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http://www.who.int/medical devices/diagnostics/Selection in-vitro diagnostics



Acronyms

ALT alanine aminotransferase
AMR antimicrobial resistance
AST aspartate aminotransferase
BMP basic metabolic panel
BUN blood urea nitrogen
CBC complete blood count

CLIA chemiluminescence immunoassay

CRP C-reactive protein
CSF cerebrospinal fluid
CVD cardiovascular disease
DST drug susceptibility testing
ECL electrochemiluminescence

World Health Organization Model List of Essential In Vitro

Diagnostics

eGFR estimated glomerular filtration rate

EIA enzyme immunoassay

ELISA enzyme linked immunosorbent assay

World Health Organization Model List of Essential

EML World rica Medicines

EPTB extrapulmonary tuberculosis
GPW WHO General Programme of Work

Hb haemoglobin HbA1c haemoglobin A1c

hCG human chorionic gonadotropin

Ht haematocrit

HTLV human T-lymphotropic virus
IGRA interferon gamma release assay
INR international normalized ratio

IVDs in vitro diagnostics

LAMP loop mediated isothermal amplification

LPA line probe assay NAT nucleic acid test

NCDs noncommunicable diseases
PQ WHO Prequalification
PT prothrombin time
RBC red blood cell count
RDT rapid diagnostic test

SAGE-IVD Strategic Advisory Group of Experts on In Vitro Diagnostics

TB Mycobacterium tuberculosis

TST tuberculin skin test
UTI urinary tract infection
VHF viral haemorrhagic fever
WBC white blood cell count
WHO World Health Organization



Preface

Introduction

The World Health Organization (WHO) published the first edition of the Model List of Essential In Vitro Diagnostics (EDL) in May 2018, in recognition that IVDs are an essential component to advance universal health coverage, address health emergencies, and promote healthier populations, which are the three strategic priorities of the WHO Thirteenth General Programme of Work (2019–2023) (GPW). The EDL is also intended to complement the WHO Model List of Essential Medicines (EML) and enhance its impact.

Objectives of the Model List of Essential In Vitro Diagnostics (EDL)

The EDL outlines a group of IVDs that are recommended by WHO for use at various levels of a tiered national health care system. The EDL is not intended to be prescriptive with respect to the IVDs listed or the levels at which such IVDs can/should be used; rather country programmes should make the ultimate decisions about which IVDs are selected and where they are implemented, based on national or regional burden of disease, unmet needs and priorities.

It is expected that the EDL will provide guidance and serve as a reference to Member States (including ministries of health, programme managers, end users such as laboratory managers, procurement officers and reimbursement systems), who are developing and/or updating lists of national essential IVDs for defining universal health coverage interventions, as well as selecting and implementing such IVDs. It will also inform United Nations agencies and nongovernmental organizations that support selection, procurement, supply, donations or provision of IVDs. Finally, it will inform and guide the medical technology private sector on IVD priorities and the IVDs needed to address global health issues.

While the EDL provides a list of important tests required at various levels of the health care system, it is important to note that the EDL itself cannot have an impact without an integrated, connected, tiered laboratory system, with adequate human resources, training, laboratory infrastructure, and regulatory/quality assurance systems. Impact also requires Member States to adopt and adapt the EDL and develop national and regional EDLs, as well as to implement the selection and supply mechanisms necessary to ensure access to the IVDs.

Scope of the first edition of the EDL

Based on the EDL selection criteria described below, the EDL consists:

- A group of general laboratory tests that can be used for routine patient care as well as for the detection and diagnosis of a wide array of disease conditions – communicable and NCDs. These IVDs are grouped by test discipline (e.g. clinical chemistry, serology, haematology, microbiology and mycology) and specific test type (e.g. bilirubin, complete blood count, etc.).
- IVDs designed for the detection, diagnosis and monitoring of each of the following WHO key disease areas: HIV, TB, malaria, HBV/HCV, and HPV and syphilis. These IVDs are grouped by disease area and analyte tested.



The EDL does *not* list specific test brands, but rather consists of IVDs described according to their biological targets. Where specific products in categories of tests contained in the EDL have been prequalified by WHO or are recommended by a WHO disease programme, a link is provided to that information, which is updated regularly.

EDL content and format

For each specific test listed in the first edition of the EDL, the following are described:

Test purpose:
 Purpose for which the test can be utilized.

• Assay format: The assay format or formats in which the test is generally

available, e.g. enzyme immunoassay, nucleic acid testing.

• Specimen type: The types of specimens that can be used for the test.

Facility level: The level of the tiered health care delivery system for which

the test is suggested, as described below.

• Link to WHO guidance: If there is existing WHO guidance available on the test or

category of testing, a link is provided to the appropriate

location on the WHO website.

• WHO PQ or endorsed products: For each specific test for which there are brands of products

either prequalified by WHO or otherwise endorsed by WHO,

a link is provided.

The EDL is presented by health care facility level in two tiers:

I IVDs for Primary health care;

II IVDs for Health care facilities with clinical laboratories.

Recommended use of the EDL

In order to effectively use the EDL and adapt it to national needs, WHO recognizes that Member States will need to consider a variety of factors. These include, among others: local demographics and burden of disease; local disease elimination priorities; local availability of treatments; training and experience of available personnel; local unmet needs and testing gaps; supply chain and transport links; quality assurance capacity; financial resources; information technology capabilities; and environmental factors.

To that end, information that supports the selection and use of IVDs on the EDL, such as relevant WHO clinical guidelines, selected systematic reviews, key references, lists of prequalified IVDs and IVDs recommended by WHO disease control departments, as well as other relevant resources on quality assurance, basic techniques, procurement and maintenance guidance, will be collated and maintained on the WHO website on an IVD-specific webpage linked to the EDL.

The EDL should not be used in isolation, but in the context of the scope of testing services that meet the clinical needs and expectations in each country through their own particular laboratory networks. An illustrative example of a tiered health care delivery and laboratory network in resource-limited countries is set out in Figure 1. The pyramid of testing reflects that there are generally a large number of primary care facilities and that they serve most patients directly for primary care needs. As one goes up the levels of the system, there are a smaller number of



centralized facilities serving fewer patients directly. In the case of national reference laboratories and some provincial laboratories, they may not serve patients directly or they may offer a broad set of specialist consultative services, and act more as referral centres for quality assurance and training or for conducting complex tests (either using samples drawn at facilities lower in the system and transported or by receiving patients referred directly from other facilities).

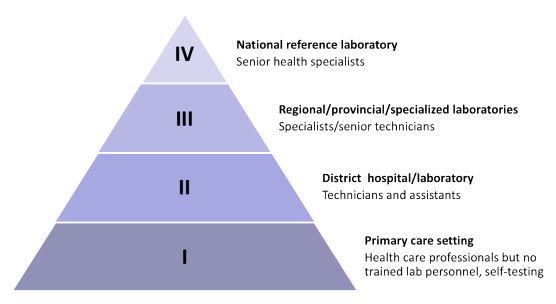


Figure 1. The types of testing that are appropriate at each tier will be country-specific and will include, among others, factors such as access to electricity, reagent grade water, phlebotomy and specialized human resources.¹

For purposes of the first edition of the EDL and to simplify its presentation and use, IVDs are listed for two tiers: primary care settings where no or minimal laboratories are available (Level I in Figure 1) or for laboratory-based facilities (Levels II, III, and IV in Figure 1).

Process of development of the first edition of the EDL

In March 2017, the WHO Expert Committee on Selection and Use of Essential Medicines recommended that an EDL be developed. In support of that recommendation, WHO created an EDL Secretariat, which drafted the first edition of the EDL in consultation with colleagues in the various WHO disease programmes. It was then posted online for open consultation. WHO also created a Strategic Advisory Group of Experts on In Vitro Diagnostics (SAGE-IVD) to support the development of the EDL and to advise on other IVD policies and initiatives. SAGE-IVD held its first meeting from 16–20 April 2018 at WHO headquarters, Geneva, where it made recommendations for the content, format and implementation of the first edition of the EDL, as well as its processes moving forward.

Selection of IVDs for inclusion in the first edition of the EDL

The selection of the diagnostics tests for the EDL took into account the following priorities:

¹ Adapted from: WHO (2017). Guidance for procurement of in vitro diagnostics and related laboratory items and equipment (http://www.who.int/diagnostics_laboratory/publications/procurement/en/).



- IVDs for primary care settings, providing an essential diagnostics package that can form the basis for screening and case management of patients at entry-level health care facilities.
- Public health approach, providing information on access to affordable, quality-assured IVDs, targeting high burden diseases, both communicable diseases and NCDs, and diseases of public health importance.
- IVDs for priority diseases such as HIV, TB, malaria, hepatitis HBV/HCV, and HPV and syphilis infections.

Specifically, the general laboratory diagnostics in the first edition of the EDL were compiled based on existing WHO guidance, guidelines and technical manuals and priority medical devices lists, which are referenced at the end of the list.

The disease-specific IVDs were selected from WHO evidence-based guidelines, which are referred to in the EDL with links to the respective documents. An additional factor considered by WHO was the availability of evidence from the WHO Prequalification of In Vitro Diagnostics Programme (PQ), or from other WHO IVD assessment processes, as applicable, which further support the choice of certain diagnostic test categories. Links to relevant documents are provided in the EDL by type of test.

Process for updating the EDL going forward

The EDL will be expanded and updated annually with the intention to ultimately cover a broad, comprehensive spectrum of disease. WHO will issue a call for applications to add IVD test categories to the next edition of the EDL in mid-2018. The call will request applicants to provide information on clinical accuracy or impact of the proposed IVDs. The first EDL will be expanded significantly over the next few years, incorporating tests for other important areas such as antimicrobial resistance, additional NCDs, emerging pathogens, emergencies and outbreaks, and neglected tropical diseases. It is foreseen that the EDL will be an important tool to increase access to appropriate, affordable, and quality-assured IVDs, particularly where they are most needed to address health priorities.

Relationship between the EDL and List of Prequalified In Vitro Diagnostics

It should be noted that the EDL and PQ List are complementary and distinct. The PQ lists include priority IVDs which have been assessed by WHO and are identified by brand (in contrast to the EDL which lists categories of IVDs). Currently the PQL has a narrower scope than the EDL.

Having IVDs on the PQ list is not a requirement for a category of tests to be considered for inclusion in the EDL. In the context of the EDL, the PQ list should be viewed as a resource as it lists specific prequalified brands of products that correspond to certain categories of tests in the EDL. Relevant links are provided in the EDL.



Implementation of the EDL by countries

It will be important that Member States adopt and adapt the EDL to develop their own national EDLs. These national EDLs will then need to be implemented to ensure impact. Implementation requires countries to invest in integrated, connected, tiered laboratory systems, with adequate human resources, training, laboratory infrastructure, and regulatory and quality assurance systems.

Glossary

Essential diagnostics: Diagnostics that satisfy the priority health care needs of the population and are selected with due regard to disease prevalence and public health relevance, evidence of efficacy and accuracy, and comparative cost-effectiveness; similar to the definition of an essential medicine.

Health care facility with laboratory support: District, regional, provincial or specialized hospitals/laboratories and national reference laboratories. Trained laboratory technicians, specialist expertise and laboratory infrastructure/equipment are available at the appropriate level. Note: All diagnostic tests available at the primary care level are assumed to be available at higher levels as appropriate.

In vitro diagnostics: A subset of medical devices, defined as: a device which, whether used alone or in combination, is intended by the manufacturer for the in vitro examination of specimens derived from the human body solely or principally to provide information for diagnostic, monitoring or compatibility purposes. It includes reagents, calibrators, control material, test kits, etc.²

Medical device: Any article, apparatus, instrument, machine, appliance, implant, reagent for in vitro use, software, material or other similar related articles, intended to be used, alone or in combination, for human beings, for one or more of the specific medical purpose(s) of:

- diagnosis, prevention, monitoring, treatment or alleviation of disease;
- diagnosis, monitoring, treatment, alleviation of or compensation for an injury;
- investigation, replacement, modification, or support of the anatomy or of a physiological process;
- supporting or sustaining life;
- control of conception;
- disinfection of medical devices:
- providing information by means of in vitro examination of specimens derived from the human body;

and does not achieve its primary intended action by pharmacological, immunological or metabolic means, in or on the human body, but which may be assisted in its intended function by such means.

Primary health care: Health centres, doctors' offices, health posts, outreach clinics. Typically, self-testing and rapid diagnostics tests are available, but there are either no laboratories, or small laboratories with trained health care personnel but no trained laboratory technicians.

² Global Harmonization Task Force (2012). Definition of the terms medical device and in vitro diagnostic (IVD) medical device (http://www.imdrf.org/docs/ghtf/final/sg1/technical-docs/ghtf-sg1-n071-2012-definition-of-terms-120516.pdf#search, accessed 3 May 2018).



Additional materials to assist countries in the selection and implementation of IVDs

These include, but are not limited to:

WHO website (www.who.int).

Guidance for procurement of in vitro diagnostics and related laboratory items and equipment. Second edition. Geneva: World Health Organization; 2017 (http://www.who.int/diagnostics_laboratory/publications/procurement/en/).

WHO Global model regulatory framework for medical devices including in vitro diagnostic medical devices. WHO Medical device technical series. Geneva: World Health Organization; 2017 (http://www.who.int/medical devices/publications/global model regulatory framework meddev/e n/).

Consultation on technical and operational recommendations for clinical laboratory testing harmonization and standardization. Geneva: World Health Organization; 2008 (http://www.who.int/healthsystems/round9 9.pdf).

Global Health Observatory data data. Geneva: World Health Organization; 2017 (http://www.who.int/gho/en/).

WHO guide for the stepwise laboratory improvement process towards accreditation in the African Region (SLIPTA). Brazzaville: WHO Regional Office for Africa; 2015.

(http://www.afro.who.int/publications/who-guide-stepwise-laboratory-improvement-process-towards-accreditation-slipta-african).

Laboratory quality standards and their implementation. WHO Regional Office for the Western Pacific and WHO Regional Office for South-East Asia; 2011

(http://www.who.int/medical devices/publications/lab quality standards/en/).

Guide for national public health laboratory networking to strengthen integrated disease surveillance and response (IDSR). Brazzaville: WHO Regional Office for Africa; 2008 (http://www.afro.who.int/publications/guide-national-public-health-laboratory-networking-strengthen-integrated-disease).

Guidance for development of national laboratory strategic plans. Brazzaville: WHO Regional Office for Africa and Atlanta (Georgia): United States Centers for Disease Control and Prevention (CDC); 2009 (http://www.who.int/hiv/amds/amds guide dev nat lab strat.pdf).

Guidance for procurement of in vitro diagnostics and related laboratory items and equipment. Geneva: World Health Organization; 2017

(http://www.who.int/diagnostics_laboratory/publications/procurement/en/).

WHO expert meeting report on short, medium and longer term product development priorities in HIV-related diagnostics. Geneva: World Health Organization; 2012 (http://www.who.int/hiv/pub/meetingreports/hiv_diagnostics/en/).

Interagency list of priority medical devices for essential interventions for reproductive, maternal, newborn and child health; 2015

(http://www.who.int/medical devices/publications/interagency med dev list/en/).



WHO list of priority medical devices for cancer management; 2017 (http://www.who.int/medical devices/publications/priority med dev cancer management/en/).

WHO publications on medical devices. Geneva: World Health Organization; 2018 (http://www.who.int/medical_devices/publications/en/).

List of Essential In Vitro Diagnostics (EDL)

The first edition of the EDL is presented by health care facility level in two tiers:

- I Primary health care; with section a for general IVDs; and section b for specific diseases.
- II Health care facilities with clinical laboratories, with section a for general IVDs; and section b for specific diseases.

I List of Essential In Vitro Diagnostics (EDL): For primary health care

Includes IVDs for health posts, community health centres, doctors' offices, outreach clinics and ambulatory care.

Typically, self-testing and rapid diagnostics tests are available, but there are either no laboratories, or only small laboratories with trained health care personnel but no trained laboratory technicians.

In case laboratory facilities are available in a primary health care facility, please refer to the IVDs described in the next tier.

It should be noted that in some cases sampling can take place where there are no laboratories, and then processed in the next tier.

I.a General IVDs	for primary health car	e		
Note: See list of	WHO supporting docui	ments at the end.		
	Diagnostic test	Test purpose	Assay format	Specimen type
Haematology	Haemoglobin (Hb)	Diagnosis and monitoring of anaemia Key clinical marker for severe infections (i.e. malaria, dengue, VHFs) Safety monitoring when using certain drugs (e.g. Zidovudine	Haemoglobinometer	Capillary whole blood Venous whole blood
		for HIV)	Dipstick	Urine
	White blood cell count	Surrogate marker for certain infections, inflammation or certain cancers (e.g. leukaemia)	Haematology analyser	Capillary whole blood Venous whole blood
	CBC manual (only as back-up to automated method)	To detect anaemia, infections and leukaemia	Haemocytometer (to measure WBC) and Wright, May-Grünwald or Giemsa stain (for differential detection of parasites, malignant cells)	Capillary whole blood Venous whole blood
			Peripheral blood film examination	Capillary whole blood Venous whole blood

I.a General IVDs for primary health care Note: See list of WHO supporting documents at the end. Diagnostic test Test purpose Assay format Specimen type Clinical chemistry and Albumin To detect/monitor kidney disease Dipstick Urine immunoassays To detect/monitor liver disease, liver/pancreas and bile duct Bilirubin Dipstick Urine disorders, and red cell destruction Glucose To diagnose and screen for diabetes and intermediate Dipstick Capillary whole blood hyperglycaemia, to diagnose hypoglycaemia Urine Capillary whole blood Glucometer Haemoglobin A1c Diagnosis and monitoring of diabetes mellitus Handheld and small analyser Capillary whole blood (HbA1c) Whole blood lactate To assess metabolic acidosis, diabetic keto-acidosis, sepsis and Electro-analytical method Arterial whole blood dehydration Handheld analyser Venous whole blood Blood transfusion To determine blood compatibility for blood transfusions; Rh Antisera for agglutination Capillary whole blood Blood typing typing for pregnant women Venous whole blood Human chorionic Serology Pregnancy Dipstick Urine gonadotropin (hCG) Microbiology, Urine dipstick and Detection of UTIs (dipstick) and identification of red and white Multi-parameter strips (dipstick) Urine mycology and urine microscopy blood cells, casts, squamous epithelial cells, bacteria, yeast, and light microscopy parasitology Schistosoma haematobium and other cellular components (microscopy) Microbial morphology, presence/absence of white blood cells Microscopy Microscopic examination of slides Disease appropriate specimens (e.g. versus squamous epithelial cells for presumptive identification as wet preparations or which venous whole blood, urine, stool, etc.) have been treated with a variety of organism-specific chemical stains (e.g. Gram stain)

	Diagnostic test	Test purpose	Assay format	Specimen type	WHO prequalified or endorsed products	WHO supporting documents
Hepatitis B	Hepatitis B surface antigen (HBsAg)	Screening for acute and chronic hepatitis B (HBV) infection: infants over 12 months of age, children, adolescents, adults	RDT	Oral fluid Capillary whole blood	http://www.who.int/di agnostics_laboratory/e valuations/pq- list/hbsag/public_repor t/en/	Guidelines on hepatitis B and C testing (February 2017): http://apps.who.int/iris/bitstream/handle/10665/25 4621/9789241549981-eng.pdf?sequence=1
	Hepatitis B e antigen (HBeAg)	Staging to assess the need for HBV treatment in chronic HBV infection	RDT	Capillary whole blood	N/A	
Hepatitis C	Antibodies to HCV (anti-HCV)	Screening for HCV infection: infants over 18 months of age, children, adolescents, adults	RDT	Oral fluid Capillary whole blood	http://www.who.int/di agnostics laboratory/e valuations/pq- list/hcv/public report/e n/	Guidelines on hepatitis B and C testing (February 2017): http://apps.who.int/iris/bitstream/handle/10665/25 4621/9789241549981-eng.pdf?sequence=1
HIV	Antibodies to HIV 1/2 (anti-HIV) test	HIV self-testing	RDT	Oral fluid Capillary whole blood	http://www.who.int/di agnostics laboratory/e valuations/pq-list/self- testing_public- report/en/	Guidelines on HIV self-testing and partner notification (2016) http://apps.who.int/iris/bitstream/handle/10665/25 1655/9789241549868-eng.pdf?sequence=1 Consolidated guidelines on HIV testing services (July 2015) http://www.who.int/hiv/pub/guidelines/hiv-
		For the diagnosis of HIV infection: adults, adolescents, children and infants over 18 months of age	RDT	Oral fluid Capillary whole blood		testing-services/en/ WHO implementation tool for pre-exposure prophylaxis (PrEP) of HIV infection, module 10 for testing providers (2017) http://www.who.int/hiv/pub/prep/prep-implementation-tool/en/

	Diagnostic test	Test purpose	Assay format	Specimen type	WHO prequalified or endorsed products	WHO supporting documents
HIV	Combined HIV antibody/p24 antigen (anti- HIV/p24 Ag) test	For the diagnosis of HIV infection: adults, adolescents, children and infants over 18 months of age	RDT	Oral fluid Capillary whole blood	http://www.who.int/di agnostics_laboratory/e valuations/pq-list/hiv- rdts/public_report/en/	Consolidated guidelines on HIV testing services (2015) http://www.who.int/hiv/pub/guidelines/hiv-testing-services/en/
Malaria	Plasmodium spp. antigens; species specific (e.g. HRP2) and/or pan-species specific (e.g. pan- pLDH)	For diagnosis of one or more human malaria species (P. falciparum, P. vivax, P. malariae, P. ovale)	RDT	Capillary whole blood	http://www.who.int/di agnostics laboratory/e valuations/pq- list/malaria/public rep ort/en/	WHO guidelines for the treatment of malaria, third edition (2015) http://apps.who.int/iris/bitstream/10665/162441/1/9789241549127 eng.pdf Malaria rapid diagnostic test performance. Results of WHO product testing of malaria RDTs: Round 7 (2015–2016) http://www.who.int/malaria/publications/atoz/978924151268/en/ WHO good practices for selecting and procuring rapid diagnostic tests for malaria (2011) http://apps.who.int/iris/bitstream/handle/10665/44530/9789241501125 eng.pdf?sequence=1
	Plasmodium spp.	For diagnosis of one or more human malaria species (P. falciparum, P. vivax, P. malariae, P. ovale and P. knowlesi) and monitoring response to treatment	Light microscopy (if good quality microscopy available)	Capillary whole blood	N/A	WHO guidelines for the treatment of malaria, third edition (2015) http://apps.who.int/iris/bitstream/10665/162441/1/9 789241549127 eng.pdf Basic malaria microscopy Part I: Learner's guide (2010) http://apps.who.int/iris/bitstream/handle/10665/442 08/9789241547826 eng.pdf?sequence=1 Malaria microscopy standard operating procedures (2015) http://www.wpro.who.int/mvp/lab_quality/mm_sop/en/

	Diagnostic test	Test purpose	Assay format	Specimen type	WHO prequalified or endorsed products (all TB tests are evaluated and	WHO supporting documents
					guidelines developed through the WHO Global TB Programme)	
Tuberculosis	Mycobacterium tuberculosis bacteria	For the diagnosis and treatment monitoring of active TB	Microscopy	Sputum	Implementing tuberculosis diagnostics: Policy framework (2015) http://apps.who.int/iris/bitstre am/10665/162712/1/9789241 508612_eng.pdf	Compendium of WHO guidelines and associated standards: Ensuring optimum delivery of the cascade of care for patients with tuberculosis (2017) http://apps.who.int/iris/bitstream/handle/10665/25918 0/9789241512572-eng.pdf?sequence=1 Implementing tuberculosis diagnostics: Policy
		For the diagnosis of active TB	LAMP	Sputum	The use of loop-mediated isothermal amplification (TB-LAMP) for the diagnosis of pulmonary tuberculosis: Policy guidance (2016) http://apps.who.int/iris/bitstream/10665/249154/1/9789241 511186-eng.pdf?ua=1	framework (2015) http://apps.who.int/iris/bitstream/10665/162712/1/978 9241508612_eng.pdf
	Immune response	For the diagnosis of latent TB infection	Intradermal skin test (TST)	N/A	Latent TB infection: Updated and consolidated guidelines for programmatic management (2018) http://apps.who.int/iris/bitstre am/handle/10665/260233/978 9241550239- eng.pdf;jsessionid=6D1BB2463 12B378ACFEBF9BFFAFEB0ED?s equence=1	

I.b Dise	Disease-specific IVDs for primary health care										
	Diagnostic test	Test purpose	Assay format	Specimen type	WHO prequalified or endorsed products	WHO supporting documents					
Syphilis	Antibodies to Treponema pallidum	For the diagnosis or as an aid in the diagnosis of <i>T.</i> pallidum	RDT	Capillary whole blood	http://www.who.int/diagnostic s_laboratory/evaluations/PQ_li st/en/	WHO laboratory diagnosis of sexually transmitted infections, including human immunodeficiency virus (2013) http://apps.who.int/iris/bitstream/handle/10665/85343/9789241505840 eng.pdf?sequence=1					
	Combined antibodies to <i>T. pallidum</i> and to HIV-1/2 (anti-HIV)	For diagnosis or as an aid in the diagnosis of HIV-1/2 and/or <i>T. pallidum</i>	RDT	Capillary whole blood	http://www.who.int/diagnostic s_laboratory/evaluations/pq- list/hiv-rdts/public_report/en/	WHO Information note on the use of dual HIV/syphilis rapid diagnostic tests (RDT) (2017) http://apps.who.int/iris/bitstream/handle/10665/252849/WHO-RHR-17.01-eng.pdf?sequence=1					

II List of Essential In Vitro Diagnostics (EDL): For health care facilities with clinical laboratories

This list includes district, regional, provincial or specialized hospitals or laboratories and national reference laboratories.

Trained laboratory technicians, specialist expertise and laboratory infrastructure/equipment are available at the appropriate level.

Note: All diagnostic tests available at the primary care level are assumed to be available at higher levels as appropriate.

The list includes: section a for general laboratory equipment; and section b tests for specific diseases.

	Diagnostic test	Test purpose	Assay format	Specimen type
Clinical chemistry and immunoassays	Alanine amino- transferase (ALT)	To assess liver function (often done with AST)	Optical and electro-analytical methods	Serum Plasma
	Albumin	To detect/monitor malnutrition, liver or kidney disease	Photometric, turbidimetric and nephelometric testing	Urine Serum Plasma
	Alkaline phosphatase	To detect/monitor malnutrition, Paget's disease or certain malignancies, including liver cancer	Colorimetric testing	Serum Plasma
	Aspartate amino- transaminase (AST)	To assess of liver function (often done with ALT)	Optical and electro-analytical methods	Serum Plasma
	Bilirubin	To detect/monitor liver disease, liver/pancreas and bile duct disorders, and red cell destruction	Optical and electro-analytical methods	Serum Plasma
	Blood pH and gases	To assess lung function, metabolic or kidney disorders, and monitor oxygen therapy Measurement of blood pH, oxygen and carbon dioxide	Electro-analytical methods, including portable analysers	Arterial whole blood Venous whole blood
	Blood urea nitrogen (BUN)	To assess kidney function and disease	Optical and electro-analytical methods	Serum Plasma
	Creatinine	To estimate glomerular filtration rate (eGFR) and urine albumin/creatinine ratio Key clinical marker for management of severe infections (i.e. sepsis, Lassa fever), as well as antimicrobial regimen adjustment	Optical and electro-analytical methods	Serum Urine
	Electrolytes	To monitor organ damage and electrolyte alterations	Optical and electro-analytical methods	Serum Plasma

II.a General IVDs for health care facilities with clinical laboratories Note: See list of WHO supporting documents at the end. Diagnostic test Assay format Specimen type Test purpose Clinical Glucose To diagnose and screen for diabetes and intermediate Automated analyser Plasma chemistry and hyperglycaemia, to diagnose hypoglycaemia Serum immunoassays Haemoglobin A1c Diagnosis and monitoring of diabetes mellitus ELISA Capillary whole blood (HbA1c) Automated analyser Venous whole blood C-reactive protein To detect inflammation as an indicator of various conditions (e.g. RDT Venous whole blood cardiovascular disease [CVD] - high sensitivity CRP required, (CRP) ΕIΑ Serum sepsis) Plasma Lipid profile To assess risk of developing CVD and type 2 diabetes by measuring Colourimetry Plasma cholesterol, triglycerides and lipoproteins Spectrophotometry Serum Basic metabolic panel Includes glucose, sodium chloride, carbon dioxide, BUN, Photometric and colourimetric testing, ion-Venous whole blood (BMP) BUN/creatinine ratio, eGFR and may include calcium selective potentiometry Serum (8-parameter automated clinical chemistry Plasma analyser) Comprehensive BMP plus magnesium, protein, albumin, globulin, alb/glob ratio, As with BMP Venous whole blood metabolic panel bilirubin (direct or total), alkaline phosphatase, ALT/AST, (14 or more parameter automated clinical Serum chemistry analyser) Plasma Amylase and lipase Colourimetric and photometric analysers To assess acute pancreatitis Serum Peritoneal fluid (Amylase) Troponin T/I For the diagnosis of myocardial infarction Enzyme immunoassay (handheld or large Venous whole blood automated instrument) Plasma Urinalysis Detection of substances in the urine associated with metabolic Automated chemical analyser Urine disorders, renal dysfunction or UTIs Blood To determine blood compatibility for blood transfusions; Rh typing Venous whole blood Blood cross-matching Antisera for agglutination transfusion for pregnant women Transfusion To screen for Chagas, HTLV in the blood supply etc. (see also EDL EIA (microplate) Serum transmitted infections sections on HIV, hepatitis C, hepatitis B, syphilis) Manual method Plasma CLIA/ECL (automated instrument) Serum Plasma Serology Human chorionic Pregnancy Optical method Serum gonadotropin (hCG)

	Diagnostic test	Test purpose	Assay format	Specimen type
Microbiology, mycology and parasitology	Urine dipstick and urine microscopy	Detection of UTIs (dipstick) and identification of red and white blood cells, casts, squamous epithelial cells, bacteria, yeast, Schistosoma haematobium and other cellular components (microscopy)	Multi-parameter strips (dipstick) and light microscopy	Urine
	Microscopy	Microbial morphology, presence/absence of white blood cells versus squamous epithelial cells for presumptive identification	Microscopic examination of slides as wet preparations or which have been treated with a variety of organism-specific chemical stains (e.g. Gram stain)	Disease appropriate specimens (e.g. venous whole blood, urine, stool, CSF, etc.)
	Culture	Initial step in the process of bacterial species detection and identification to support selection of appropriate antibiotic treatment regimens	Culture on growth media plates and incubator followed by recovery of isolates and species identification (traditional manual techniques or automated equipment)	Disease appropriate specimens (e.g. venous whole blood, urine, stool, CSF etc.)
	Blood culture	For the diagnosis of bacterial and fungal blood stream infections (sepsis)	Blood culture bottle and incubator followed by recovery of isolates and species identification (traditional manual techniques or automated equipment)	Venous whole blood
	Antimicrobial susceptibility testing	Final step in the process of selection of appropriate antibiotic treatment regimens after species identification	Antimicrobial susceptibility testing of isolates – may be done manually using disc diffusion technique or using automated platforms	Microbial isolates
Haematology	Haematocrit (Ht)	Diagnosis and monitoring of anaemia Volume of red blood cells as a percentage of total blood volume	Microhaematocrit centrifuge	Capillary or venous whole blood
	Prothrombin time test and international normalized ratio (PT/INR)	To detect/diagnose a bleeding disorder or excessive clotting disorder (PT); monitor performance of anticoagulant medications (INR)	Handheld or automated coagulation analyser	Citrate plasma
	Platelet count	Diagnosis of thrombocytopenia	Haemocytometer	Capillary whole blood
		Marker to manage severe infections associated with bleeding and sepsis (i.e. VHF, meningococcemia) and certain haematological disorders	Haematology analyser	Venous whole blood
	Complete blood count (CBC) Automated, differential	Evaluation of patient's overall health and to detect a wide range of disorders, including anaemia, infection and leukaemia	Automated hematology analyser (WBC, RBC, platelets, Hb and Ht) includes lymphocytes, monocytes and granulocytes (for three-part differential)	Venous whole blood

II.b Disease-specific IVDs for health care facilities with clinical laboratories WHO prequalified or endorsed Diagnostic test **Assay format** Specimen type WHO supporting documents Test purpose products Hepatitis B Hepatitis B surface Screening for acute and RDT Venous whole blood http://www.who.int/diagnostic Guidelines on hepatitis B and C testing s laboratory/evaluations/pgantigen (HBsAg) chronic hepatitis B (HBV) Plasma (February 2017) infection: infants over 12 list/hbsag/public report/en/ http://apps.who.int/iris/bitstream/handle/ Serum 10665/254621/9789241549981months of age, children, EIA Plasma eng.pdf?sequence=1 adolescents, adults Serum CLIA Plasma Serum Virological Staging to assess the NAT Serum (HBV DNA need for HBV treatment Plasma in chronic HBV infection quantitative) and monitoring of response to treatment Staging to assess the ΕIΑ N/A Hepatitis B e antigen Serum (HBeAg) need for HBV treatment Plasma in chronic HBV infection CLIA N/A Serum Plasma IgM-specific For the diagnosis of acute EIA (microplate) N/A Serum antibodies to HBV infection – used for Manual method Plasma hepatitis B core outbreak investigation CLIA/ECL Serum N/A antigen (IgM anti-(automated Plasma HBc) instrument) Antibodies to Determining EIA (microplate) Serum N/A hepatitis B surface effectiveness of HBV Manual method Plasma antigen (anti-HBs) immunization at patient and at a population level Also used as a marker for CLIA/ECL N/A Serum recovery from HBV (automated Plasma infection instrument)

	Diagnostic test	Test purpose	Assay format	Specimen type	WHO prequalified or endorsed products	WHO supporting documents
Hepatitis C	Antibodies to HCV (anti-HCV)	Screening for HCV infection: infants over 18 months of age, children,	RDT	Venous whole blood Plasma Serum	http://www.who.int/diagnostic s laboratory/evaluations/pq- list/hcv/public report/en/	Guidelines on hepatitis B and C testing (February 2017) http://apps.who.int/iris/bitstream/h
		adolescents, adults	EIA (microplate) Manual method	Serum Plasma		andle/10665/254621/978924154998 1-eng.pdf?sequence=1
			CLIA/ECL (automated instrument)	Serum Plasma		
	Antibodies to HCV (anti-HCV) and HCV	Screening for HCV past or present infection: infants	EIA (microplate) Manual method	Serum Plasma		
	core antigen (HCV cAg)	over 18 months of age, children, adolescents, adults	CLIA/ECL (automated instrument)	Serum Plasma		
	HCV core antigen (HCV cAg)	For the diagnosis of viraemic HCV infection	CLIA/ECL (automated instrument)	Serum Plasma		
	HCV RNA (qualitative or quantitative)	For the diagnosis of viraemic HCV infection and monitoring of response to treatment as a test of cure	NAT	Serum Plasma		

	Diagnostic test	Test purpose	Assay format	Specimen type	WHO prequalified or endorsed products	WHO supporting documents
HIV	Antibodies to HIV-1/2 (anti-HIV) test	For the diagnosis of HIV infection: adults, adolescents, children and infants over 18 months of age	RDT	Venous whole blood Plasma Serum	http://www.who.int/diagnostic s_laboratory/evaluations/pq- list/self-testing_public- report/en/	Guidelines on HIV self-testing and partner notification (2016) http://apps.who.int/iris/bitstream/handle/10665/251655/9789241549868-eng.pdf?sequence=1
			EIA (microplate) Manual method	Serum Plasma		Consolidated guidelines on HIV testing services (July 2015)
			CLIA/ECL (automated instrument)	Serum Plasma		http://www.who.int/hiv/pub/guidelines/hiv-testing-services/en/
			,			WHO implementation tool for pre- exposure prophylaxis (PrEP) of HIV infection, module 10 for testing providers (2017) <a href="http://www.who.int/hiv/pub/prep/prep-
implementation-tool/en/">http://www.who.int/hiv/pub/prep/prep- implementation-tool/en/
		For screening for HIV in the blood supply and in blood products	EIA (microplate) Manual method CLIA/ECL	Serum Plasma Serum	N/A	Screening donated blood for transfusion transmissible infections: Recommendations (2009)
		'	(automated instrument)	Plasma		http://apps.who.int/iris/bitstream/handle/ 10665/44202/9789241547888_eng.pdf?se quence=1&isAllowed=y
	Combined HIV antibody/p24 antigen (anti-HIV/p24 Ag) test	For the diagnosis of HIV infection: adults, adolescents, children and	RDT	Venous whole blood Plasma Serum	http://www.who.int/diagnostic s laboratory/evaluations/pq- list/hiv-rdts/public report/en/	Consolidated guidelines on HIV testing services (2015) http://apps.who.int/iris/bitstream/handle/
		infants over 18 months of age	EIA (microplate) Manual method	Serum Plasma		10665/179870/9789241508926 eng.pdf?s equence=1
			CLIA/ECL (automated instrument)	Serum Plasma		
		For screening for HIV in the blood supply and in	EIA (microplate) Manual method	Serum Plasma	N/A	Screening donated blood for transfusion transmissible infections: Recommendation
		blood products	CLIA/ECL (automated instrument)	Serum Plasma		(2009)http://apps.who.int/iris/bitstream/handle/10665/44202/9789241547888_eng.pdf?sequence=1&isAllowed=y

	Diagnostic test	Test purpose	Assay format	Specimen type	WHO prequalified or endorsed products	WHO supporting documents
HIV	HIV qualitative virological or quantitative virological	For the diagnosis of HIV infection in infants under 18 months of age	NAT	Capillary whole blood Venous whole blood Dried blood spot Serum Plasma	http://www.who.int/diagnostic s_laboratory/evaluations/pq- list/hiv-vrl/public_report/en/	Consolidated guidelines on the use of antiretroviral drugs for treating and preventing HIV infection (2016) http://www.who.int/hiv/pub/arv/arv-2016/en/
	HIV quantitative virological	Monitoring of response to antiviral treatment	NAT	Dried blood spot Serum Plasma	http://www.who.int/diagnostic s_laboratory/evaluations/pq- list/hiv-vrl/public_report/en/	
	CD4 cell enumeration (quantitative)	For staging of advanced HIV disease	Flow cytometry	Capillary whole blood Venous whole blood	http://www.who.int/diagnostic s_laboratory/evaluations/pq- list/hiv-vrl/public_report/en/	
	Cryptococcal antigen test	For screening and diagnosis of cryptococcal meningitis in people living with advanced HIV	RDT	CSF Venous whole blood Serum Plasma	N/A	Guidelines for the diagnosis, prevention, and management of cryptococcal disease in HIV-infected adults, adolescents and children (2018)
		disease	EIA	CSF Serum Plasma		http://apps.who.int/iris/bitstream/handl 10665/260399/9789241550277- eng.pdf?sequence=1

	Diagnostic test	Test purpose	Assay format	Specimen type	WHO prequalified or endorsed products	WHO supporting documents
1alaria	Plasmodium spp. antigens; species specific (e.g. HRP2) and/or pan-species specific (e.g. pan- pLDH)	For diagnosis of one or more human malaria species (<i>P. falciparum, P. vivax, P. malariae, P. ovale</i>)	RDT	Capillary whole blood Venous whole blood	http://www.who.int/diagnostic s_laboratory/evaluations/pq- list/malaria/public_report/en/	WHO guidelines for the treatment of malaria, third edition (2015) http://apps.who.int/iris/bitstream/10669 162441/1/9789241549127_eng.pdf Malaria rapid diagnostic test performance Results of WHO product testing of malaria RDTs: Round 7 (2015–2016) http://www.who.int/malaria/publicationatoz/978924151268/en/ WHO good practices for selecting and procuring rapid diagnostic tests for malaria (2011) http://apps.who.int/iris/bitstream/handla 10665/44530/9789241501125_eng.pdf?equence=1
	Plasmodium spp.	For diagnosis of one or more human malaria species (<i>P. falciparum, P. vivax, P. malariae, P. ovale</i> and <i>P. knowlesi</i>) and monitoring response to treatment	Light microscopy	Capillary whole blood Venous whole blood	N/A	WHO guidelines for the treatment of malaria, third edition (2015) http://apps.who.int/iris/bitstream/1066! 162441/1/9789241549127_eng.pdf Basic malaria microscopy Part I: Learner's guide (2010) http://apps.who.int/iris/bitstream/handl 10665/44208/9789241547826_eng.pdf? quence=1 Malaria microscopy standard operating procedures (2015) http://www.wpro.who.int/mvp/lab_qual/mm_sop/en/

	Diagnostic test	Test purpose	Assay format	Specimen type	WHO prequalified or endorsed products	WHO supporting documents
Malaria	Glucose-6-phosphate dehydrogenase activity (G6PD)	To determine G6PD activity (normal, intermediate, deficient) and specifically to inform decision to administer 8-aminoquinoline group drugs for radical cure of <i>P. vivax</i> For screening newborns for G6PD deficiency	Semi quantitative fluorescent spot test	Venous whole blood	http://www.who.int/diagnostic s_laboratory/evaluations/pq- list/malaria/public_report/en/	Beutler E, Blume KG, Kaplan JC, Lohr GW, Ramot B, Valentine WN. International Committee for Standardization in Haematology: Recommended screening test for glucose-6-phosphate dehydrogenase deficiency. Br J Haematol 1979;43:469–477 WHO guidelines for the treatment of malaria, third edition (2015) http://apps.who.int/iris/bitstream/10665162441/1/9789241549127_eng.pdf

	Diagnostic test	Test purpose	Assay format	Specimen type	WHO prequalified or endorsed products (all TB tests are evaluated and guidelines developed through the WHO Global TB Programme)	WHO supporting documents
Tuberculosis	Mycobacterium tuberculosis bacteria	For the diagnosis and treatment monitoring of active TB	Microscopy	Other specimen types	Implementing tuberculosis diagnostics: Policy framework (2015) http://apps.who.int/iris/bitstream/10665/162712/1/9789241508612 eng.pdf	Compendium of WHO guidelines and associated standards: Ensuring optimum delivery of the cascade of care for patients with tuberculosis (2017)
		For the diagnosis and treatment monitoring of active TB including drug- resistant TB	Bacterial culture	Sputum or other specimen types		http://apps.who.int/iris/bitstream/hand/10665/259180/9789241512572-eng.pdf?sequence=1 Implementing tuberculosis diagnostics:
	M. tuberculosis DNA	For the diagnosis of active TB and simultaneous detection of rifampicin resistance	Cartridge-based NAT	Sputum or EPTB specimen types	WHO Meeting report of a technical expert consultation: Non-inferiority analysis of Xpert MTB/RIF Ultra compared to Xpert MTB/RIF (2017) http://apps.who.int/iris/bitstream/handle/10665/2 54792/WHO-HTM-TB-2017.04-eng.pdf;jsessionid=E02D0994930EDBD9A4BC5BB3D 3A28568?sequence=1 Automated real-time nucleic acid amplification technology for rapid and simultaneous detection of tuberculosis and rifampicin resistance: Policy update (2013) http://apps.who.int/iris/bitstream/10665/112472/1 /9789241506335 eng.pdf	Policy framework (2015) http://apps.who.int/iris/bitstream/10669 /162712/1/9789241508612 eng.pdf
	M. tuberculosis DNA mutations associated with resistance	For the detection of resistance for first- line anti-TB medicines	Molecular LPA	Sputum	The use of molecular line probe assays for the detection of resistance to isoniazid and rifampicin: Policy update (2016) http://apps.who.int/iris/bitstream/10665/250586/1/9789241511261-eng.pdf?ua=1	
	M. tuberculosis DNA mutations associated with resistance	For the detection of resistance for second-line anti-TB medicines	Molecular LPA	Sputum	The use of molecular line probe assays for the detection of resistance to second-line antituberculosis drugs: Policy update (2016) http://apps.who.int/iris/bitstream/handle/10665/246131/9789241510561-eng.pdf?sequence=1	

	Diagnostic test	Test purpose	Assay format	Specimen type	WHO prequalified or endorsed products (all TB tests are evaluated and guidelines developed through the WHO Global TB Programme)	WHO supporting documents
Tuberculosis	M. tuberculosis culture-based DST	To detect resistance to first-line and/or second-line anti-TB medicines	DST	Bacterial culture of <i>M.</i> tuberculosis	Technical report on critical concentrations for drug susceptibility testing of medicines used in the treatment of drug-resistant tuberculosis (2018) http://www.who.int/tb/publications/2018/WHO te chnical report concentrations TB drug susceptibil ity/en/	
	Lipoarabino- mannan (LAM) antigen	To aid in the diagnosis of TB in seriously ill HIV- positive inpatients	RDT	Urine	The use of lateral flow urine lipoarabinomannan assay (LF-LAM) for the diagnosis and screening of active tuberculosis in people living with HIV: Policy update (2015) http://apps.who.int/iris/bitstream/handle/10665/193633/9789241509633 eng.pdf;jsessionid=9A9EB8 86DC17658BF7FDF86758D7A9F9?sequence=1	
	Immune response	For the diagnosis of latent TB infection	IGRA	Venous whole blood	Latent TB Infection: Updated and consolidated guidelines for programmatic management (2018) http://apps.who.int/iris/bitstream/handle/10665/2 60233/9789241550239-eng.pdf;jsessionid=6D1BB246312B378ACFEBF9BFFA FEB0ED?sequence=1	

	Diagnostic test	Test purpose	Assay format	Specimen type	WHO prequalified or endorsed products	WHO supporting documents
HPV	Human papillomavirus (HPV) DNA	For cervical cancer screening	Nucleic acid test	Cervical cells collected in test specific transport fluid	http://www.who.int/diagnostics_laboratory/ evaluations/pq-list/public_report_hpv/en/	WHO human papillomavirus laboratory manual, first edition (2009) http://apps.who.int/iris/bitstream/handle/10665/70505/WHO_IVB_10.12_eng.pdf?sequence=1
Syphilis	Antibodies to Treponema pallidum	For diagnosis or as an aid in the diagnosis of <i>T. pallidum</i>	RDT EIA (Microplate)	Venous whole blood Plasma Serum Serum	http://www.who.int/diagnostics_laboratory/ evaluations/PQ_list/en/	WHO laboratory diagnosis of sexually transmitted infections, including human immunodeficiency virus (2013) http://apps.who.int/iris/bitstream/hand
			Manual method CLIA/ECL (automated instrument)	Plasma Serum Plasma		e/10665/85343/9789241505840 eng.pd f?sequence=1
		For screening blood and blood products	EIA (Microplate) Manual method	Serum Plasma	N/A	Screening donated blood for transfusion transmissible infections (2009) http://apps.who.int/iris/bitstream/handle/10665/44202/9789241547888 eng.pd f?sequence=1&isAllowed=y
	Combined antibodies to <i>T. pallidum</i> and to HIV-1/2 (anti-HIV)	For the diagnosis or as an aid in the diagnosis of HIV-1/2 and/or <i>T. pallidum</i>	RDT	Venous whole blood Plasma Serum	http://www.who.int/diagnostics_laboratory/ evaluations/pq-list/hiv- rdts/public_report/en/	WHO Information note on the use of dual HIV/syphilis rapid diagnostic tests (RDT) (2017) http://apps.who.int/iris/bitstream/handle/10665/252849/WHO-RHR-17.01-eng.pdf?sequence=1

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