

EN
ANNEX I
GENERAL COMMON SPECIFICATIONS

Part I – Requirements for performance characteristics of devices covered by Annexes II to XIII

Performance characteristics	Requirement
All performance characteristics set out in Section 9.1, points (a) and (b), Section 9.3 and Section 9.4, point (a), of Annex I to Regulation (EU) 2017/746	<ol style="list-style-type: none"> 1. The determination of performance characteristics shall be carried out in direct comparison with a state-of-the-art device. The device used for comparison shall be one bearing CE marking, if on the market at the time of the performance evaluation. 2. Devices used for determination of status of samples used in determination of performance characteristics shall be state-of-the-art devices bearing CE marking. 3. If discrepant results are identified as part of determination of performance characteristics, these results shall be resolved as far as possible, by one or more of the following: <ul style="list-style-type: none"> – by evaluation of the discrepant sample in further devices, – by use of an alternative method or marker, – by a review of the clinical status and diagnosis of the patient, – by the testing of follow-up samples. 4. The determination of performance characteristics shall be performed on a population equivalent to the European population.
Whole system failure rate	<ol style="list-style-type: none"> 5. As part of the required risk analysis the whole system failure rate leading to false negative results shall be determined in repeat assays on low-positive specimens.

Analytical sensitivity and analytical specificity, interference	6. For devices intended for use with plasma the manufacturer shall verify the performance of the device using all anticoagulants which the manufacturer indicates for use with the device, for at least 50 plasma specimens (for devices intended for detection and/or quantification of infectious agents, 25 positive and 25 negative).
Analytical and diagnostic specificity, interference and cross-reactivity	7. The manufacturer shall select the potential interfering substances to be evaluated taking account of the composition of the reagents and configuration of the device.
Batch-to-batch consistency	<p>8. For devices intended to detect antigens and antibodies, the manufacturer's batch testing criteria shall ensure that every batch consistently identifies the relevant antigens, epitopes, and antibodies and is suitable for the claimed specimen types.</p> <p>9. The manufacturer's batch release testing for first-line assays, except those covered by Tables 1 and 2 of Annex XIV, shall include at least 100 specimens negative for the relevant analyte.</p>

Part II – Requirements for performance characteristics of devices referred to in Annexes III to XIII

Performance characteristic	Requirement
Analytical and diagnostic sensitivity	<ol style="list-style-type: none"> 10. Devices intended by the manufacturer for testing body fluids other than serum or plasma, e.g. urine, saliva, etc., shall meet the same requirements as serum or plasma devices. The manufacturer shall test samples from the same individuals in both the devices to be approved and in a respective serum or plasma device. 11. Devices for self-testing shall meet the same requirements as respective devices for professional use. 12. Positive specimens used in the performance evaluation shall be selected to reflect different stages of the respective disease(s), different antibody patterns, different genotypes, different subtypes, mutants, etc. 13. Seroconversion panels shall start with a negative bleed(s) and shall reflect narrow bleeding intervals as far as possible. Where this is not possible, manufacturers shall provide a justification in the performance evaluation report. 14. For devices intended by the manufacturer to be used with serum and plasma the performance evaluation must demonstrate serum to plasma equivalency. This shall be demonstrated for at least 25 positive donations. 15. For devices detecting or quantifying antigens or nucleic acids, the target antigen or gene respectively shall be specified in the instructions for use. 16. For devices detecting or quantifying antibodies against an infectious agent, the target antigen(s) of those antibodies shall be specified in the instructions for use.
Analytical and diagnostic specificity	<ol style="list-style-type: none"> 17. Devices intended by the manufacturer for testing body fluids other than serum or plasma, e.g. urine, saliva, etc., shall meet the same requirements as serum or plasma devices. The performance evaluation shall test samples from the same individuals in both the devices to be approved and in a respective serum or plasma device. 18. Devices for self-testing shall meet the same requirements as respective devices for professional use. 19. Negative specimens used in a performance evaluation shall be defined so as to reflect the target population for which the device is intended, such as blood donors, hospitalised patients, pregnant women, etc. 20. Specificity shall be calculated using the frequency of repeatedly reactive (i.e. false positive) results in samples negative for the target marker.

	21. For devices intended by the manufacturer to be used with serum and plasma the performance evaluation must demonstrate serum to plasma equivalency. This shall be demonstrated for at least 25 negative donations.
Analytical and diagnostic specificity, interference and cross-reactivity	22. The manufacturer shall include specimens such as, where applicable: <ul style="list-style-type: none"> – specimens representing related infections, – specimens from multipara, i.e. women who have had more than one pregnancy, or rheumatoid factor (RF) positive patients, – specimens containing human antibodies to components of the expression system, for example anti-<i>E. coli</i>, or anti-yeast.
Performances obtained by lay persons	23. Relevant parts of the performance evaluation shall be carried out (or repeated) by appropriate lay persons to validate the operation of the device and the instructions for use. The lay persons selected for the performance evaluation shall be representative of the intended users groups.

ANNEX II

COMMON SPECIFICATIONS FOR DEVICES INTENDED FOR DETECTION OF BLOOD GROUP ANTIGENS IN THE ABO, RH, KELL, DUFFY AND KIDD BLOOD GROUP SYSTEMS

Scope

This Annex applies to devices intended for detection of blood group antigens in the ABO, Rh, Kell, Duffy and Kidd blood group systems.

Table 1 applies to performance evaluation of devices detecting blood group antigens in the ABO, Rh, Kell, Duffy and Kidd blood group systems. Table 2 applies to manufacturer's batch-to-batch consistency testing of reagents and reagent products to determine blood group antigens in the ABO, Rh, Kell, Duffy and Kidd blood group systems (test reagents, control materials).

Table 1. Performance evaluation of devices detecting blood group antigens in the ABO, Rh, Kell, Duffy and Kidd blood group systems

Reagent specificity	Number of tests per method claimed by the manufacturer	Total number of samples to be tested for a launch device	Total number of samples to be tested for a new formulation, or use of well-characterised reagents	General qualification criteria	Specific qualification criteria	Acceptance criteria
Anti-ABO1 (Anti-A), Anti-ABO2 (Anti-B), Anti-ABO3 (Anti-A,B)	≥500	≥3 000	≥1 000	Clinical samples: 10 % of the test population Neonatal specimens: > 2 % of the test population	ABO samples shall include > 40 % A and B antigen positive samples which may include samples from group A, group B and group AB	All of the reagents shall show comparable performance to state-of-the-art CE marked devices with regard to claimed reactivity of the device. For CE marked devices where the application or use has been changed or extended, further testing shall be carried out in accordance with the
Anti-RH1 (Anti-D)	≥500	≥3 000	≥1 000		Performance evaluation of Anti-D reagents shall include tests against a	

					range of weak RH1 (D) and partial RH1 (D) samples, depending on the intended use of the product. Weak and/or partial D cells shall account for > 2 % of RH1 (D) positive samples.	requirements outlined in column 2 above (“Number of tests per method claimed by the manufacturer”).
Anti-RH2 (Anti-C), Anti-RH4 (Anti-c), Anti- RH3 (Anti-E)	≥100	≥1 000	≥200			
Anti-RH5 (Anti-e)	≥100	≥500	≥200			
Anti-KEL1 (Anti-K)	≥100	≥500	≥200			
Anti-JK1 (Jk ^a), Anti-JK2 (Jk ^b)	≥100	≥500	≥200			
Anti-FY1 (Fy ^a), Anti-FY2 (Fy ^b)	≥100	≥500	≥200			

Note: Positive specimens used in the performance evaluation shall be selected to reflect variant and weak antigen expression.

Table 2. Manufacturer’s batch-to-batch consistency testing of reagents and reagent products to determine blood group antigens in the ABO, Rh, Kell, Duffy and Kidd blood group systems

1. Test reagents

Blood group reagents	Minimum number of control cells to be tested as part of specificity testing					Acceptance criteria		
	Positive reactions					Negative reactions		
	A1	A2B	Ax			B	O	
Anti-ABO1(Anti-A)	2	2	2 ¹			2	2	
	B	A1B				A1	O	
Anti-ABO2(Anti-B)	2	2				2	2	
	A1	A2	Ax	B		O		
Anti-ABO3(Anti-A,B)	2	2	2 ¹	2		4		
	R1r	R2r	WeakD			r'r	r''r	rr
Anti-RH1 (Anti-D)	2	2	2 ¹			1	1	1
	R1R2	R1r	r'r			R2R2	r''r	rr
Anti-RH2 (Anti-C)	2	1	1			1	1	1
	R1R2	R1r	r'r			R1R1		
Anti-RH4 (Anti-c)	1	2	1			3		
	R1R2	R2r	r''r			R1R1	r'r	rr
Anti-RH3 (Anti-E)	2	1	1			1	1	1
	R1R2	R2r	r''r			R2R2		
Anti-RH5 (Anti-e)	2	1	1			3		
	Kk					kk		
Anti-KEL1 (Anti-K)	4					3		
	Jk(a+b+)					Jk(a-b+)		
Anti-JK1 (Anti-Jk ^a)	4					3		

Each batch of reagent shall exhibit unequivocal positive or negative results by all techniques claimed by the manufacturer in accordance with the results obtained from the performance evaluation data.

	Jk(a+b+)					Jk(a+b-)			
Anti-JK2 (Anti-Jk ^b)	4					3			
	Fy(a+b+)					Fy(a-b+)			
Anti-FY1 (Anti-Fy ^a)	4					3			
	Fy(a+b+)					Fy(a+b-)			
Anti-FY2 (Anti-Fy ^b)	4					3			

Note: Polyclonal reagents shall be tested against a wider panel of cells to confirm specificity and exclude presence of unwanted contaminating antibodies.

¹ Only where reactivity against these antigens is claimed.

2. Control materials (red cells)

The phenotype of red cells used in the control of blood typing reagents listed above shall be confirmed using established device.

ANNEX III

COMMON SPECIFICATIONS FOR DEVICES INTENDED FOR DETECTION OR QUANTIFICATION OF MARKERS OF HUMAN IMMUNODEFICIENCY VIRUS (HIV) INFECTION

Scope

1. This Annex applies to devices intended for detection or quantification of markers of human immunodeficiency virus (HIV) infection.

Table 1 applies to first-line assays for HIV-1/2 antibody (anti-HIV 1/2) and first-line combined antigen/antibody assays for HIV 1/2 (HIV 1/2 Ag/Ab) which are not rapid tests.

Table 2 applies to first-line assays for anti-HIV 1/2 and HIV 1/2 Ag/Ab which are rapid tests.

Table 3 applies to confirmatory assays for anti-HIV 1/2.

Table 4 applies to antigen tests for HIV-1 and HIV Ag/Ab assays.

Table 5 applies to for qualitative and quantitative NAT devices for HIV ribonucleic acid (RNA).

Table 6 applies to HIV-1/2 self-tests.

Definitions

2. For the purposes of this Annex, the following definitions apply:

- (1) 'seroconversion HIV sample' means:
 - p24 antigen and/or HIV RNA positive, and
 - recognised by the antibody first-line assays, and
 - positive or indeterminate in confirmatory assays.
- (2) 'early seroconversion HIV sample' means:
 - p24 antigen and/or HIV RNA positive, and
 - not recognised by the antibody first-line assays, and

- indeterminate or negative in confirmatory assays.

Table 1. First-line assays: anti-HIV 1/2, HIV 1/2 Ag/Ab (requirements for antibody detection)

Performance characteristic	Specimen	Specimen numbers, features, use	Acceptance criteria
Diagnostic sensitivity	Positive specimens	<p>≥400 HIV-1 ≥100 HIV-2 including 40 non-B-subtypes including 25 positive ‘same day’ fresh serum samples (≤ 1 day after sampling)</p> <p>all available HIV/1 subtypes shall be represented by at least 3 samples per subtype</p>	all true positive samples shall be identified as positive
	Seroconversion panels	<p>≥20 panels ≥10 further panels (at notified body or manufacturer) at least 40 early seroconversion HIV samples shall be tested</p>	diagnostic sensitivity during seroconversion shall represent the state of the art all seroconversion HIV samples shall be identified as positive
Diagnostic specificity	Unselected blood donors (including first-time donors) ¹	≥5 000	≥99,5%
	Hospitalised patients	≥200	Potential limitations for specificity, if any, shall be identified
Cross-reactivity	Potentially cross-reacting specimens	<p>≥100 in total (such as RF+, from related virus infections, from pregnant women, subjects recently vaccinated against any infectious agent)</p>	

¹ Blood donor populations shall be investigated from at least two blood donation centres and consist of consecutive blood donations, which have not been selected to exclude first-time donors.

Table 2. Rapid tests: anti-HIV 1/2, HIV 1/2 Ag/Ab (requirements for antibody detection)

Performance characteristic	Specimen	Specimen numbers, features, use, acceptance criteria	Acceptance criteria
Diagnostic sensitivity	Positive specimens	≥ 400 HIV-1 ≥ 100 HIV-2 including 40 non-B-subtypes all available HIV/1 subtypes shall be represented by at least 3 samples per subtype	all true positive samples shall be identified as positive
	Seroconversion panels	≥ 20 panels ≥ 10 further panels (at notified body or manufacturer) at least 40 early seroconversion HIV samples shall be tested	diagnostic sensitivity during seroconversion shall represent the state of the art all seroconversion HIV samples shall be identified as positive
Diagnostic specificity	Unselected blood donors (including first-time donors)	$\geq 1\ 000$	$\geq 99\ %$
	Hospitalised patients	≥ 200	Potential limitations for specificity, if any, shall be identified
Cross-reactivity	Potentially cross-reacting specimens	≥ 200 samples from pregnant women ≥ 100 other potentially cross-reacting specimens in total (e.g. RF+, from related infections)	

Table 3. Confirmatory assays: anti-HIV 1/2

Performance characteristic	Specimen	Specimen numbers, features, use	Acceptance criteria

Diagnostic sensitivity	Positive specimens	≥ 200 HIV-1 ≥ 100 HIV-2 Including different stages of infection and reflecting different antibody patterns	Identification as “confirmed positive” or “indeterminate”, not as “negative”
	Seroconversion panels	≥ 15 seroconversion panels/low titre panels ≥ 40 early seroconversion HIV samples shall be tested	Diagnostic sensitivity during seroconversion shall represent the state of the art All seroconversion HIV samples shall be identified as positive
Diagnostic specificity	Blood donors	≥ 200	No false positive results / no neutralisation
	Hospitalised patients	≥ 200	
Cross-reactivity	Potentially cross-reacting specimens	≥ 50 in total (including samples from pregnant women, samples with indeterminate results in other confirmatory assays)	

Table 4. Antigen tests: HIV-1, HIV Ag/Ab (requirements for antigen detection)

Performance characteristic	Specimen	Specimen numbers, features, use	Acceptance criteria
Diagnostic sensitivity	Positive specimens	≥ 50 HIV-1 antigen positive ≥ 50 cell culture supernatants including different HIV-1 subtypes and HIV-2	all true positive samples shall be identified as positive (after neutralisation if applicable)
	Seroconversion panels	≥ 20 seroconversion panels/low titre panels ≥ 40 early seroconversion HIV samples as referred to in Section 2(2) of this Annex shall be tested	diagnostic sensitivity during seroconversion shall represent the state of the art all seroconversion HIV samples shall be identified as positive

Analytical sensitivity	First International Reference Reagent HIV-1 p24 Antigen, NIBSC code: 90/636		≤ 2 IU/ml
Diagnostic specificity	Blood donors	≥200	≥ 99,5 % after neutralisation or, if no neutralisation test available, after resolution of the sample status
	Hospitalised patients	≥200	
Cross-reactivity	Potentially cross-reacting specimens	≥50	Potential limitations for specificity, if any, shall be identified

Table 5. Qualitative and quantitative NAT devices for HIV RNA

1. For target sequence amplification devices, a functionality control for each specimen (internal control) shall reflect the state of the art. This control shall as far as possible be used throughout the whole process, i.e. extraction, amplification/hybridisation, detection.
2. Genotype and/or subtype detection shall be demonstrated by appropriate primer or probe design validation and shall also be validated by testing characterised genotyped samples.
3. Potential cross-reactivity of non-target nucleic acid sequences shall be analysed by appropriate primer or probe design validation and shall also be validated by testing selected samples
4. Results of quantitative NAT devices shall be traceable to international standards or calibrated reference materials, if available, and be expressed in international units utilised in the specific field of application.
5. Qualitative HIV NAT devices intended to be used to detect the presence of HIV in blood, blood components, cells, tissues or organs, or in any of their derivatives, in order to assess their suitability for transfusion, transplantation or cell administration shall be designed to detect both HIV-1 and HIV-2.
6. Qualitative HIV NAT devices, other than virus typing devices, shall be designed to compensate for the potential failure of a HIV-1 NAT target region by using two independent target regions.

Performance characteristic	Specimen	Specimen numbers, features, use	Acceptance criteria
Analytical sensitivity	WHO International Standard HIV-1 RNA; WHO International Standard HIV-2 RNA; or calibrated reference materials	NAT sensitivity and NAT LOD shall be validated by dilution series of reference materials, testing of replicates (minimal 24) at different analyte concentrations, including those with transition from positive to negative results in the respective NAT assay. LOD shall be expressed as 95% positive cut-off value (IU/ml) after statistical analysis (e.g. Probit). ¹ quantitative NAT: definition of lower, upper quantification limit, precision, accuracy, 'linear' measuring range, 'dynamic range'. Reproducibility at different concentration levels	According to the state of the art
HIV geno-/subtype sensitivity	all relevant genotypes/subtypes, preferably from international reference materials potential substitutes for rare HIV subtypes (to be quantified by appropriate methods): cell culture supernatants; in vitro transcripts; plasmids	Qualitative NAT: at least 10 samples/genotype or subtype Quantitative NAT: dilution series for demonstration of quantification efficiencies	According to the state of the art
Diagnostic sensitivity	Positive specimens reflecting the routine conditions of users (e.g. no pre-selection of specimens)	Quantitative NAT: ≥ 100 Comparative results with another NAT test system shall be generated in parallel	According to the state of the art
	Seroconversion panels	Qualitative NAT: ≥ 10 panels Comparative results with another NAT test system shall be generated in parallel	According to the state of the art
Diagnostic specificity	Blood donor specimens	Qualitative NAT: ≥ 500 Quantitative NAT: ≥ 100	According to the state of the art
Cross-reactivity	Potentially cross-reacting specimens	≥ 10 human retrovirus positive samples (e.g. HTLV)	

Carry-over	High HIV RNA positive; HIV RNA negative	At least five runs with alternating high-positive and negative specimens shall be performed during robustness studies. The virus titres of the high positive samples shall be representative of high virus titres occurring naturally.	According to the state of the art
Detection in relation to antibody status	HIV-RNA positives: anti-HIV negative, anti-HIV positive	Pre-seroconversion (anti-HIV negative) and post-seroconversion (anti-HIV positive) specimens	According to the state of the art
Whole system failure rate	HIV RNA low positive	≥100 HIV RNA low-positive specimens shall be tested. These specimens shall contain a virus concentration equivalent to three times the 95 % positive cut-off virus concentration.	≥99% positive

¹ Reference: European Pharmacopeia 9.0, 2.6.21 Nucleic acid amplification techniques, Validation.

Table 6. Additional requirements for HIV-1/2 self-tests

Performance characteristic	Specimens¹	Specimen number
Result interpretation supervised ²	Interpretation of tests carried out with contrived specimens ³ by lay users reflecting a range of results: <ul style="list-style-type: none"> • non-reactive • reactive • weak reactive⁴ • invalid 	≥ 100
Diagnostic sensitivity	lay users that are known positive	≥ 200
Diagnostic specificity	lay users that do not know their status	≥ 400

	Lay users that are at high risk of acquiring the infection	≥ 200
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¹ For each body fluid claimed for use with the device, e.g. whole blood, urine, saliva, etc. The sensitivity and specificity of the device for self-testing in the hands of lay users shall be defined against the confirmed patient infectious status.

² The result interpretation study shall include reading and interpretation of the contrived test results by 100 lay people with each lay person subjected to read one or more contrived tests from the specified range of result reactivity levels. The manufacturer shall determine the concordance between lay user reading and professional user reading.

³ Tests shall be performed prior to the result interpretation study using whenever possible the specimen type intended by the manufacturer.

⁴ A higher proportion of the specimens shall be in the weak-positive range close to the cut-off or LOD of the test.

ANNEX IV

COMMON SPECIFICATIONS FOR DEVICES INTENDED FOR DETECTION OR QUANTIFICATION OF MARKERS OF HUMAN T-CELL LYMPHOTROPIC VIRUS (HTLV) INFECTION

Scope

This Annex applies to devices intended for detection or quantification of markers of human T-cell lymphotropic virus (HTLV) infection.

Table 1 applies to first-line assays for antibodies against HTLV I or II (anti-HTLV I/II) which are not rapid tests.

Table 2 applies to first-line assays for anti HTLV I/II which are rapid tests.

Table 3 applies to confirmatory assays for anti-HTLV I/II.

Table 4 applies to NAT devices for HTLV I/II.

Table 1. First-line assays: anti-HTLV I/II

Performance characteristic	Specimen	Specimen numbers, features, use	Acceptance criteria
Diagnostic sensitivity	Positive specimens	≥ 300 HTLV-I ≥ 100 HTLV-II including 25 positive ‘same day’ fresh serum samples (≤ 1 day after sampling)	all true positive samples shall be identified as positive
	Seroconversion panels	To be defined when available	diagnostic sensitivity during seroconversion shall represent the state of the art, if applicable
Diagnostic specificity	Unselected blood donors (including first-time donors) ¹	$\geq 5\ 000$	$\geq 99,5\%$
	Hospitalised patients	≥ 200	Potential limitations for specificity, if any, shall be identified
Cross-reactivity	Potentially cross-reacting specimens	≥ 100 in total (e.g. RF+, from related virus infections, from pregnant women,)	

¹ Blood donor populations shall be investigated from at least two blood donation centres and consist of consecutive blood donations, which have not been selected to exclude first-time donors.

Table 2. Rapid tests: anti HTLV I/II

Performance characteristic	Specimen	Specimen numbers, features, use, acceptance criteria	Acceptance criteria
Diagnostic sensitivity	Positive specimens	≥300 HTLV-I ≥100 HTLV-II	all true positive samples shall be identified as positive
	Seroconversion panels	To be defined when available	diagnostic sensitivity during seroconversion shall represent the state of the art, if applicable
Diagnostic specificity	Unselected blood donors (including first-time donors)	≥1000	≥ 99%
	Hospitalised patients	≥200	Potential limitations for specificity, if any, shall be identified
Cross-reactivity	Potentially cross-reacting specimens	≥200 samples from pregnant women ≥100 other potentially cross-reacting specimens in total (e.g. RF+, from related infections)	

Table 3. Confirmatory assays: anti-HTLV I/II

Performance characteristic	Specimen	Specimen numbers, features, use, acceptance criteria	Acceptance criteria
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Diagnostic sensitivity	Positive specimens	≥200 HTLV I ≥100 HTLV II	Identification as “confirmed positive” or “indeterminate”, not as “negative”
	Seroconversion panels	To be defined when available	diagnostic sensitivity during seroconversion shall represent the state of the art, if applicable
Diagnostic specificity	Blood donors	≥200	No false positive results
	Hospitalised patients	≥200	
Cross-reactivity	Potentially cross-reacting specimens	≥50 in total (including samples from pregnant women, samples with indeterminate results in other confirmatory assays)	

Table 4. NAT devices for HTLV I/II

1. For target sequence amplification devices, a functionality control for each specimen (internal control) shall reflect the state of the art. This control shall as far as possible be used throughout the whole process, i.e. extraction, amplification/hybridisation, detection.
2. Genotype and/or subtype detection shall be demonstrated by appropriate primer or probe design validation and shall also be validated by testing characterised genotyped samples.
3. Potential cross-reactivity of non-target nucleic acid sequences shall be analysed by appropriate primer or probe design validation and shall also be validated by testing selected samples.
4. Results of quantitative NAT devices shall be traceable to international standards or calibrated reference materials, if available, and be expressed in international units utilised in the specific field of application.

Performance characteristic	Specimen	Specimen numbers, features, use, acceptance criteria	Acceptance criteria
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Analytical sensitivity	International reference preparations	NAT sensitivity and NAT LOD shall be validated by dilution series of reference materials, testing of replicates (minimal 24) at different analyte concentrations, including those with transition from positive to negative results in the respective NAT assay. LOD shall be expressed as 95% positive cut-off value (IU/ml) after statistical analysis (e.g. Probit). ¹ quantitative NAT: definition of lower, upper quantification limit, precision, accuracy, 'linear' measuring range, 'dynamic range'. Reproducibility at different concentration levels	According to the state of the art
HTLV I and HTLV II genotype sensitivity	all relevant genotypes, preferably from international reference materials potential substitutes for rare HTLV genotypes (to be quantified by appropriate methods): cell culture supernatants; in vitro transcripts; plasmids	Qualitative NAT: at least 10 samples/genotype or subtype Quantitative NAT: dilution series for demonstration of quantification efficiencies	According to the state of the art
Diagnostic specificity	Blood donor specimens	Qualitative NAT: ≥ 500 Quantitative NAT: ≥ 100	According to the state of the art
Cross-reactivity	Potentially cross-reacting specimens	≥ 10 human retrovirus positive samples (e.g. HIV-1, HIV-2)	According to the state of the art
Carry-over	High HTLV RNA positive; HTLV RNA negative	At least five runs with alternating high-positive and negative specimens shall be performed during robustness studies. The virus titres of the high positive samples shall be representative of high virus titres occurring naturally.	According to the state of the art

Detection in relation to antibody status	HTLV-RNA positives: anti-HTLV negative, anti-HTLV positive	Pre-seroconversion (anti-HTLV negative) and post-seroconversion (anti-HTLV positive) specimens	According to the state of the art
Whole system failure rate	HTLV RNA low positive	≥100 HTLV RNA low-positive specimens shall be tested. These specimens shall contain a virus concentration equivalent to three times the 95 % positive cut-off virus concentration.	≥99% positive

¹ Reference: European Pharmacopeia 9.0, 2.6.21 Nucleic acid amplification techniques, Validation.

ANNEX V

COMMON SPECIFICATIONS FOR DEVICES INTENDED FOR DETECTION OR QUANTIFICATION OF MARKERS OF HEPATITIS C VIRUS (HCV) INFECTION

Scope

This Annex applies to devices intended for detection or quantification of markers of hepatitis C virus (HCV) infection.

Table 1 applies to first-line assays for anti-HCV antibodies (anti-HCV) and combined antigen/antibody tests for HCV (HCV Ag/Ab) which are not rapid tests.

Table 2 applies to first-line assays for anti-HCV and HCV Ag/Ab which are rapid tests.

Table 3 applies to confirmatory and supplemental assays for anti-HCV.

Table 4 applies to HCV antigen tests and HCV Ag/Ab.

Table 5 applies to qualitative and quantitative NAT devices for HCV RNA.

Table 6 applies to HCV self-tests.

Table 7 applies to HCV serotyping and genotyping assays.

Table 1. First-line assays: anti-HCV, HCV Ag/Ab (requirements for antibody detection)

Performance characteristic	Specimen	Specimen numbers, features, use	Acceptance criteria
Diagnostic sensitivity	Positive specimens	≥ 400 Including samples from different stages of infection and reflecting different antibody patterns HCV genotype 1-4: > 20 samples per genotype (including non-a subtypes of genotype 4); HCV genotypes 5 and 6: > 5 samples each; including 25 positive 'same day' fresh serum samples (≤ 1 day after sampling)	all true positive samples shall be identified as positive

	Seroconversion panels	<p>≥20 panels ≥10 further panels (at notified body or manufacturer)</p> <p>HCV seroconversion panels for the evaluation of HCV antigen and antibody combined tests (HCV Ag/Ab) shall start with one or more negative bleeds and comprise panel members from early HCV infection (HCV core antigen and/or HCV RNA positive but anti-HCV negative).</p>	<p>diagnostic sensitivity during seroconversion shall represent the state of the art</p> <p>HCV Ag/Ab tests shall demonstrate enhanced sensitivity in early HCV infection when compared to HCV antibody only tests.</p>
Diagnostic specificity	Unselected blood donors (including first-time donors) ¹	≥5 000	≥99,5%
	Hospitalised patients	≥200	Potential limitations for specificity, if any, shall be identified
Cross-reactivity	Potentially cross-reacting specimens	<p>≥100 in total (e.g. RF+, from related virus infections, from pregnant women,)</p>	

¹ Blood donor populations shall be investigated from at least two blood donation centres and consist of consecutive blood donations, which have not been selected to exclude first-time donors.

Table 2. Rapid tests: anti-HCV, HCV Ag/Ab (requirements for antibody detection)

Performance characteristic	Specimen	Specimen numbers, features, use, acceptance criteria	Acceptance criteria
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Diagnostic sensitivity	Positive specimens	<p>≥400</p> <p>including samples from different stages of infection and reflecting different antibody patterns.</p> <p>HCV genotype 1-4: > 20 samples per genotype (including non-a subtypes of genotype 4); HCV genotypes 5 and 6: > 5 samples each;</p>	all true positive samples shall be identified as positive
	Seroconversion panels	<p>≥20 panels</p> <p>≥10 further panels (at notified body or manufacturer)</p> <p>HCV seroconversion panels for the evaluation of HCV antigen and antibody combined tests (HCV Ag/Ab) shall start with one or more negative bleeds and comprise panel members from early HCV infection (HCV core antigen and/or HCV RNA positive but anti-HCV negative).</p>	<p>diagnostic sensitivity during seroconversion shall represent the state of the art</p> <p>HCV Ag/Ab tests shall demonstrate enhanced sensitivity in early HCV infection when compared to HCV antibody only tests.</p>
Diagnostic specificity	Unselected blood donors (including first-time donors) ¹	≥1 000	≥ 99 %
	Hospitalised patients	≥200	Potential limitations for specificity, if any, shall be identified
Cross-reactivity	Potentially cross-reacting specimens	<p>≥200 samples from pregnant women</p> <p>≥100 other cross-reacting specimens in total (e.g. RF+, from related infections)</p>	

Table 3. Confirmatory and supplemental assays: anti-HCV

Performance characteristic	Specimen	Specimen numbers, features, use, acceptance criteria	Acceptance criteria
Diagnostic sensitivity	Positive specimens	<p>≥300</p> <p>Including specimens from different stages of infection and reflecting different antibody patterns.</p>	identification as “confirmed positive” or “indeterminate”, not as “negative”

		HCV genotypes 1 – 4: > 20 specimens (including non-a subtypes of genotype 4; HCV genotype 5: > 5 specimens; HCV genotype 6: as far as available	
	Seroconversion panels	≥15 seroconversion panels/low titre panels	diagnostic sensitivity during seroconversion shall represent the state of the art
Diagnostic specificity	Blood donors	≥200	No false positive results/ no neutralisation
	Hospitalised patients	≥200	
Cross-reactivity	Potentially cross-reacting specimens	≥50 in total (including samples from pregnant women, samples with indeterminate results in other confirmatory assays)	

Table 4. Antigen tests: HCV antigen, HCV Ag/Ab (requirements for antigen detection)

Performance characteristic	Specimen	Specimen numbers, features, use, acceptance criteria	Acceptance criteria
Diagnostic sensitivity	Positive specimens	≥25 HCV core antigen and/or HCV RNA positive but anti-HCV negative specimens, comprising HCV genotypes 1-6 (if a genotype is not available, a justification shall be made)	all true positive specimens shall be identified as positive
	Seroconversion panels	≥20 seroconversion panels/low titre panels HCV seroconversion panels for the evaluation of HCV antigen and antibody combined tests shall start with one or more negative bleeds and comprise panel members from early HCV infection (HCV core antigen and/or HCV RNA positive but anti-HCV negative).	diagnostic sensitivity during seroconversion shall represent the state of the art HCV antigen and antibody combined tests shall demonstrate enhanced sensitivity in early HCV infection when compared to HCV antibody only tests.
Analytical sensitivity	WHO International Standard HCV core (PEI 129096/12)	Dilution series	

Diagnostic specificity	Blood donors	≥200	≥ 99,5 % after neutralisation or, if no neutralisation test available, after resolution of the sample status
	Hospitalised patients	≥200	Potential limitations for specificity, if any, shall be identified
Cross-reactivity	Potentially cross-reacting specimens	≥50	

Table 5. Qualitative and quantitative NAT devices for HCV RNA

1. For target sequence amplification devices, a functionality control for each specimen (internal control) shall reflect the state of the art. This control shall as far as possible be used throughout the whole process, i.e. extraction, amplification/hybridisation, detection.
2. Genotype and/or subtype detection shall be demonstrated by appropriate primer or probe design validation and shall also be validated by testing characterised genotyped samples.
3. Potential cross-reactivity of non-target nucleic acid sequences shall be analysed by appropriate primer or probe design validation and shall also be validated by testing selected samples.
4. Results of quantitative NAT devices shall be traceable to international standards or calibrated reference materials, if available, and be expressed in international units utilised in the specific field of application.

Performance characteristic	Specimen	Specimen numbers, features, use, acceptance criteria	Acceptance criteria
Analytical sensitivity	WHO International Standard HCV RNA(or calibrated reference materials)	NAT sensitivity and NAT LOD shall be validated by dilution series of reference materials, testing of replicates (minimal 24) at different analyte concentrations, including those with transition from positive to negative results in the respective NAT assay. LOD shall be expressed as 95% positive cut-off value (IU/ml) after statistical analysis (e.g. Probit). ¹ quantitative NAT: definition of lower, upper quantification limit, precision, accuracy, 'linear' measuring range, 'dynamic range'. Reproducibility at different concentration levels	According to the state of the art

HCV genotype sensitivity	all relevant genotypes/subtypes, preferably from international reference materials potential substitutes for rare HCV genotypes (to be quantified by appropriate methods): in vitro transcripts; plasmids	Qualitative NAT: ≥ 10 samples/genotype or subtype Quantitative NAT: dilution series for demonstration of quantification efficiencies	According to the state of the art
Diagnostic sensitivity	Positive specimens reflecting the routine conditions of users (e.g. no pre-selection of specimens)	Quantitative NAT: ≥ 100 Comparative results with another NAT test system shall be generated in parallel	According to the state of the art
	Seroconversion panels	Qualitative NAT: ≥ 10 panels Comparative results with another NAT test system shall be generated in parallel	According to the state of the art
Diagnostic specificity	Blood donor specimens	Qualitative NAT: ≥ 500 Quantitative NAT: ≥ 100	According to the state of the art
Cross-reactivity	Potentially cross-reacting specimens	>10 human flavivirus (e.g. HGV, YFV) positive samples	According to the state of the art
Carry-over	High HCV RNA positive; HCV RNA negative	At least five runs with alternating high-positive and negative specimens shall be performed during robustness studies. The virus titres of the high positive samples shall be representative of high virus titres occurring naturally.	According to the state of the art
Detection in relation to antibody status	HCV RNA positives: anti-HCV negative, anti-HCV positive	Pre-seroconversion (anti-HCV negative) and post-seroconversion (anti-HCV positive) specimens	According to the state of the art
Whole system failure rate	HCV RNA low positive	≥ 100 HCV RNA low-positive specimens shall be tested. These specimens shall contain a virus concentration equivalent to three times the 95 % positive cut-off virus concentration.	$\geq 99\%$ positive

¹ Reference: European Pharmacopeia 9.0, 2.6.21 Nucleic acid amplification techniques, Validation.

Table 6. Additional requirements for HCV self-tests

Performance characteristic	Specimens¹	Number of specimens
Result interpretation supervised ²	Interpretation of tests carried out with contrived specimens ³ by lay users reflecting a range of results: <ul style="list-style-type: none"> • non-reactive • reactive • weak reactive⁴ • invalid 	≥ 100
Diagnostic sensitivity	lay users that are known positive	≥ 200
Diagnostic specificity	lay users that do not know their status	≥ 400
	lay users that are at high risk of acquiring the infection	≥ 200

¹ For each body fluid claimed for use with the device, e.g. whole blood, urine, saliva, etc. The sensitivity and specificity of the device for self-testing in the hands of lay users shall be defined against the confirmed patient infectious status.

² The result interpretation study shall include reading and interpretation of the contrived test results by 100 lay people with each lay person subjected to read one or more contrived tests from the specified range of result reactivity levels. The manufacturer shall determine the concordance between lay user reading and professional user reading.

³ Tests shall be performed prior to the result interpretation study using whenever possible the specimen type intended by the manufacturer.

⁴ A higher proportion of the specimens shall be in the weak-positive range close to the cut-off or LOD of the test.

Table 7. HCV serotyping and genotyping assays

1. For target sequence amplification devices, a functionality control for each specimen (internal control) shall reflect the state of the art. This control shall as far as possible be used throughout the whole process, i.e. extraction, amplification/hybridisation, detection.

2. Genotype and/or subtype detection shall be demonstrated by appropriate primer or probe design validation and shall also be validated by testing characterised genotyped samples.
3. Potential cross-reactivity of non-target nucleic acid sequences shall be analysed by appropriate primer or probe design validation and shall also be validated by testing selected samples.
4. Results of quantitative NAT devices shall be traceable to international standards or calibrated reference materials, if available, and be expressed in international units utilised in the specific field of application.

Performance characteristic	Specimen	Specimen numbers, features, use, acceptance criteria	Acceptance criteria
Diagnostic sensitivity	Positive specimens	≥ 200 Including specimens from different stages of infection and reflecting different antibody patterns HCV genotype 1-6: > 20 specimens per genotype	$\geq 95\%$ agreement between serotyping and genotyping; $> 95\%$ agreement between genotyping and sequencing
Diagnostic specificity	Negative specimens	≥ 100	

ANNEX VI

COMMON SPECIFICATIONS FOR DEVICES INTENDED FOR DETECTION OR QUANTIFICATION OF MARKERS OF HEPATITIS B VIRUS (HBV) INFECTION

Scope

This Annex applies to devices intended for detection or quantification of markers of hepatitis B virus (HBV) infection.

Table 1 applies to first-line assays for hepatitis B surface antigen (HBsAg), and for antibodies against hepatitis B core antigen (anti-HBc) which are not rapid tests.

Table 2 applies to first-line assays for HBsAg and anti-HBc which are rapid tests.

Table 3 applies to confirmatory assays for HBsAg.

Table 4 applies to assays for the hepatitis B virus markers: hepatitis B surface antibodies (anti-HBs), IgM antibody against the hepatitis B core antigen (anti-HBc IgM), antibodies against the hepatitis Be antigen (anti-HBe) and hepatitis Be antigen (HBeAg).

Table 5 applies to qualitative and quantitative NAT for HBV deoxyribonucleic acid (DNA).

Table 6 applies to HBV self-tests.

Table 1. First-line assays: HBsAg, anti-HBc

Performance characteristic	Specimen	Specimen numbers, features, use	Acceptance criteria
Diagnostic sensitivity	Positive specimens	≥400 anti-HBc: including evaluation of different HBV markers HBsAg: including different HBV genotypes / subtypes / mutants anti-HBc or HBsAg: including 25 positive 'same day' fresh serum (≤ 1 day after sampling)	Overall performance shall be at least equivalent to that of the state of the art device referred to in Annex I (3).

	Seroconversion panels	HBsAg assays: ≥20 panels ≥10 further panels (at notified body or manufacturer) antiHBc assays: to be defined when available	diagnostic sensitivity during seroconversion shall represent the state of the art (for antiHBc this shall be the case if applicable)
Analytical sensitivity	WHO Third International Standard HBsAg (subtypes ayw1/adw2, HBV genotype B4, NIBSC code: 12/226)		For HBsAg assays: <0,130 IU/ml
Diagnostic specificity	Unselected blood donors (including first-time donors) ¹	≥5 000	≥99,5%
	Hospitalised patients	≥200	Potential limitations for specificity, if any, shall be identified
Cross-reactivity	Potentially cross-reacting specimens	≥100 in total (e.g. RF+, from related virus infections, from pregnant women,)	

¹Blood donor populations shall be investigated from at least two blood donation centres and consist of consecutive blood donations, which have not been selected to exclude first-time donor.

Table 2. Rapid tests: HBsAg, anti-HBc

Performance characteristic	Specimen	Specimen numbers, features, use	Acceptance criteria
Diagnostic sensitivity	Positive specimens	≥400 including evaluation of different HBV markers including different HBV genotypes / subtypes / mutants	Overall performance shall be at least equivalent to that of state of the art device referred to in Annex I (3)

	Seroconversion panels	HBsAg assays: ≥20 panels ≥10 further panels (at notified body or manufacturer) Anti-HBc assays: to be defined when available	diagnostic sensitivity during seroconversion shall represent the state of the art (for anti-HBc this shall be the case if applicable)
Diagnostic specificity	Unselected blood donors (including first-time donors)	≥1 000	HBsAg assays: ≥ 99 % anti-HBc assays: ≥ 99 %
	Hospitalised patients	≥200	Potential limitations for specificity, if any, shall be identified
Cross-reactivity	Potentially cross-reacting specimens	≥200 samples from pregnant women ≥100 other potentially cross-reacting specimens in total (e.g. RF+, from related infections)	

Table 3. Confirmatory assays: HBsAg

Performance characteristic	Specimen	Specimen numbers, features, use, acceptance criteria	Acceptance criteria
Diagnostic sensitivity	Positive specimens	≥300 Including samples from different stages of infection Including 20 'high positive' samples (>26 IU/ml); 20 samples in the cut-off range	Correct identification as positive (or indeterminate), not negative
	Seroconversion panels	≥15 seroconversion panels/low titre panels	diagnostic sensitivity during seroconversion shall represent the state of the art
Analytical sensitivity	Third International Standard for HBsAg, subtypes ayw1/adw2, HBV genotype B4, NIBSC code: 12/226		
Diagnostic specificity	Negative specimens	≥10 false positives as available from the performance evaluation of the first-line assay	No false positive results/ no neutralisation

Cross-reactivity	Potentially cross-reacting specimens	≥50	
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Table 4. Assays for the HBV markers: anti-HBs, anti-HBc IgM, anti-HBe, HBeAg

Performance characteristic		Anti-HBs	Anti-HBc IgM	Anti-HBe	HBeAg	Acceptance criteria
Diagnostic sensitivity	Positive specimens	≥100 vaccinees ≥100 naturally infected persons	≥200 Including samples from different stages of infection (acute/chronic, etc.)	≥200 Including samples from different stages of infection (acute/chronic, etc.)	≥200 Including samples from different stages of infection (acute/chronic, etc.)	≥ 98 % (for Anti-HBc IgM: applicable only on samples from acute infection stage)
	Seroconversion panels	10 follow-ups or anti-HBs seroconversions	When available	When available	When available	Diagnostic sensitivity during seroconversion shall represent the state of the art (for Anti-HBc IgM, Anti-HBe, HBeAg this shall be the case if applicable)
Analytical sensitivity	Standards	WHO 2nd International Standard for anti-hepatitis B surface antigen (anti-HBs) immunoglobulin, human NIBSC code: 07/164		WHO 1st International Standard Anti-hepatitis B virus e antigen (anti-HBe), PEI code 129095/12	WHO 1st International Standard for Hepatitis B Virus e Antigen (HBeAg) PEI code 129097/12 HBe	Anti-HBs: < 10 mIU/ml
Diagnostic specificity	Negative specimens	≥500 Including clinical samples ≥50 potentially interfering samples	≥200 blood donations ≥200 clinical samples ≥50 potentially interfering samples	≥200 blood donations ≥200 clinical samples ≥50 potentially interfering samples	≥200 blood donations ≥200 clinical samples ≥50 potentially interfering samples	≥ 98 %

Table 5. Qualitative and quantitative NAT devices for HBV DNA

1. For target sequence amplification devices, a functionality control for each specimen (internal control) shall reflect the state of the art. This control shall as far as possible be used throughout the whole process, i.e. extraction, amplification/hybridisation, detection.
2. Genotype and/or subtype detection shall be demonstrated by appropriate primer or probe design validation and shall also be validated by testing characterised genotyped samples.
3. Potential cross-reactivity of non-target nucleic acid sequences shall be analysed by appropriate primer or probe design validation and shall also be validated by testing selected samples.
4. Results of quantitative NAT devices shall be traceable to international standards or calibrated reference materials, if available, and be expressed in international units utilised in the specific field of application.

Performance characteristic	Specimen	Specimen numbers, features, use	Acceptance criteria
Analytical sensitivity	WHO International Standard HBV DNA (or calibrated reference materials)	NAT sensitivity and NAT LOD shall be validated by dilution series of reference materials, testing of replicates (minimal 24) at different analyte concentrations, including those with transition from positive to negative results in the respective NAT assay. LOD shall be expressed as 95% positive cut-off value (IU/ml) after statistical analysis (e.g. Probit). ¹ quantitative NAT: definition of lower, upper quantification limit, precision, accuracy, 'linear' measuring range, 'dynamic range'. Reproducibility at different concentration levels	According to the state of the art
HBV genotype sensitivity	WHO International Reference Panel HBV DNA (HBV genotypes) all relevant genotypes/subtypes, preferably from international reference materials	Qualitative NAT: at least 10 samples/genotype or subtype Quantitative NAT: dilution series for demonstration of quantification efficiencies	According to the state of the art

	potential substitutes for rare HBV genotypes (to be quantified by appropriate methods): plasmids; synthetic DNA		
Diagnostic sensitivity	Positive specimens reflecting the routine conditions of users (e.g. no pre-selection of specimens)	Quantitative NAT: ≥ 100 Comparative results with another NAT test system shall be generated in parallel	According to the state of the art
	Seroconversion panels	Qualitative NAT: ≥ 10 panels Comparative results with another NAT test system shall be generated in parallel	According to the state of the art
Diagnostic specificity	Blood donor specimens	Qualitative NAT: ≥ 500 Quantitative NAT: ≥ 100	According to the state of the art
Cross-reactivity	Potentially cross-reacting specimens		According to the state of the art
Carry-over	High HBV DNA positive; HBV DNA negative	At least five runs with alternating high-positive and negative specimens shall be performed during robustness studies. The high positive samples shall comprise samples with naturally occurring high virus titres.	According to the state of the art
Detection in relation to antibody status	HBV DNA positives: anti-HBV negative, anti-HBV positive	Pre-seroconversion (anti-HBV negative) and post-seroconversion (anti-HBV positive) specimens	According to the state of the art
Whole system failure rate	HBV DNA low positive	≥ 100 HBV DNA low-positive specimens shall be tested. These specimens shall contain a virus concentration equivalent to three times the 95 % positive cut-off virus concentration.	$\geq 99\%$ positive

¹ Reference: European Pharmacopeia 9.0, 2.6.21 Nucleic acid amplification techniques, Validation.

Table 6. Additional requirements for HBV self-tests

Performance characteristic	Specimens ¹	Numbers of specimens
Result interpretation supervised ²	Interpretation of tests carried out with contrived specimens ³ by lay users reflecting a range of results: <ul style="list-style-type: none"> • non-reactive • reactive • weak reactive⁴ • invalid 	≥100
Diagnostic sensitivity	lay users that are known positive	≥200
Diagnostic specificity	lay users that do not know their status	≥400
	lay users that are at high risk of acquiring the infection	≥200

¹ For each body fluid claimed for use with the device, e.g. whole blood, urine, saliva, etc. The sensitivity and specificity of the device for self-testing in the hands of lay users shall be defined against the confirmed patient infectious status.

² The result interpretation study shall include reading and interpretation of the contrived test results by 100 lay people with each lay person subjected to read one or more contrived tests from the specified range of result reactivity levels. The manufacturer shall determine the concordance between lay user reading and professional user reading.

³ Tests shall be performed prior to the result interpretation study using whenever possible the specimen type intended by the manufacturer.

⁴ A higher proportion of the specimens shall be in the weak-positive range close to the cut-off or LOD of the test.

ANNEX VIII

COMMON SPECIFICATIONS FOR DEVICES INTENDED FOR DETECTION OR QUANTIFICATION OF MARKERS OF HEPATITIS D VIRUS (HDV) INFECTION

Scope

This Annex applies to devices intended for detection or quantification of markers of hepatitis D virus (HDV) infection.

Table 1 applies to devices intended for detection (including confirmation) or quantification of the following hepatitis D virus markers: antibodies against hepatitis D virus (anti-HDV), IgM antibodies against hepatitis D virus (anti-HDV IgM), the delta antigen.

Table 2 applies to qualitative and quantitative NAT for HDV RNA.

Table 1. Assays for HDV markers: anti-HDV, anti-HDV IgM, delta antigen

Performance characteristic		Anti-HDV	Anti-HDV IgM	Delta antigen	Acceptance criteria
Diagnostic sensitivity	Positive specimens	≥100 Specifying markers of HBV coinfection	≥50 Specifying markers of HBV coinfection	≥10 Specifying markers of HBV coinfection	≥ 98 %
Diagnostic specificity	Negative specimens	≥200 Including clinical samples ≥50 potentially interfering samples	≥200 Including clinical samples ≥50 potentially interfering samples	≥200 Including clinical samples ≥50 potentially interfering samples	≥ 98 %

Table 2. Qualitative and quantitative NAT devices for HDV RNA

1. For target sequence amplification devices, a functionality control for each specimen (internal control) shall reflect the state of the art. This control shall as far as possible be used throughout the whole process, i.e. extraction, amplification/hybridisation, detection.
2. Genotype and/or subtype detection shall be demonstrated by appropriate primer or probe design validation and shall also be validated by testing characterised genotyped samples.

3. Potential cross-reactivity of non-target nucleic acid sequences shall be analysed by appropriate primer or probe design validation and shall also be validated by testing selected samples.
4. Results of quantitative NAT devices shall be traceable to international standards or calibrated reference materials, if available, and be expressed in international units utilised in the specific field of application.

Performance characteristic	Specimen	Specimen numbers, features, use	Acceptance criteria
Analytical sensitivity	1st WHO International Standard HDV RNA, PEI code 7657/12	NAT sensitivity and NAT LOD shall be validated by dilution series of reference materials, testing of replicates (minimal 24) at different analyte concentrations, including those with transition from positive to negative results in the respective NAT assay. LOD shall be expressed as 95% positive cut-off value (IU/ml) after statistical analysis (e.g. Probit). ¹ quantitative NAT: definition of lower, upper quantification limit, precision, accuracy, 'linear' measuring range, 'dynamic range'. Reproducibility at different concentration levels	According to the state of the art
HDV genotype sensitivity	all relevant genotypes/subtypes, preferably from international reference materials potential substitutes for rare HDV genotypes (to be quantified by appropriate methods): plasmids; synthetic RNA	Quantitative NAT: dilution series for demonstration of quantification efficiencies	According to the state of the art
Diagnostic specificity	Blood donor specimens	Qualitative NAT: ≥ 100 Quantitative NAT: ≥ 100	According to the state of the art
Cross-reactivity	Potentially cross-reacting specimens		According to the state of the art

Carry-over	High HDV RNA positive; HDV RNA negative	At least five runs with alternating high-positive and negative specimens shall be performed during robustness studies. The high positive samples shall comprise samples with naturally occurring high virus titres.	According to the state of the art
Whole system failure rate	HDV RNA low positive	≥100 HDV RNA low-positive specimens shall be tested. These specimens shall contain a virus concentration equivalent to three times the 95 % positive cut-off virus concentration.	≥99% positive

¹Reference: European Pharmacopeia 9.0, 2.6.21 Nucleic acid amplification techniques, Validation.

ANNEX VIII

COMMON SPECIFICATIONS FOR DEVICES INTENDED FOR DETECTION OF MARKERS OF VARIANT CREUTZFELDT-JACOB (vCJD) DISEASE

Scope

This Annex applies to devices intended for detection of markers of variant Creutzfeldt-Jakob disease (vCJD).

Table 1 applies to devices intended for detection of markers of vCJD.

Table 1. Devices for detection of markers of vCJD

Performance characteristic	Material	Number of specimens	Acceptance criteria
Analytical sensitivity	vCJD brain spikes in human plasma (WHO reference number NHBY0/0003)	≥24 replicates of each of three dilutions of the material WHO number NHBY0/0003 (1×10^4 , 1×10^5 , 1×10^6)	23 of the 24 replicates detected at 1×10^4
	vCJD spleen spikes in human plasma (10% spleen homogenate — NIBSC reference number NHSY0/0009)	≥24 replicates of each of three dilutions of the material NIBSC number NHSY0/0009 (1×10 , 1×10^2 , 1×10^3)	23 of the 24 replicates detected at 1×10
Diagnostic sensitivity	Specimens from appropriate animal models	As many specimens as reasonably possible and available, and ≥10 specimens	90%
	Specimens from humans with known clinical vCJD	As many specimens as reasonably possible and available, and ≥10 specimens	90%
		Only in cases where 10 specimens are not available: — the number of specimens tested shall be between 6 and 9 — all available specimens shall be tested	max. one false negative result

Analytical specificity	Potentially cross-reacting blood-specimens	≥ 100	
Diagnostic specificity	Normal human plasma samples from area of low bovine spongiform encephalopathy (BSE) exposure	≥ 5000	$\geq 99,5 \%$

ANNEX IX

COMMON SPECIFICATIONS FOR DEVICES INTENDED FOR DETECTION OR QUANTIFICATION OF MARKERS OF CYTOMEGALOVIRUS (CMV) INFECTION

Scope

This Annex applies to devices intended for detection or quantification of markers of cytomegalovirus (CMV) infection.

Table 1 applies to first-line assays for total antibodies against CMV (total anti-CMV) and IgG antibodies against CMV (anti-CMV IgG).

Table 2 applies to qualitative and quantitative NAT devices for CMV DNA.

Table 1. First-line assays: total anti-CMV and anti-CMV IgG

Performance characteristic	Specimens	Total anti-CMV, anti-CMV IgG	Acceptance criteria
Diagnostic sensitivity	Positive specimens	≥400 including specimens from recent and past CMV infection, low and high positive titre samples	≥ 99% sensitivity for confirmable past infection ¹ ; overall sensitivity including recent infection ² shall be comparable to other CE-marked tests
	Seroconversion panels	To be tested when available	Diagnostic sensitivity during seroconversion shall represent the state of the art
Analytical sensitivity	Standards	WHO international standard anti-CMV IgG (PEI-code 136616/17) In case of titre determinations and quantitative statements	
Diagnostic specificity	Negative specimens	≥400 ³ CMV negatives from unselected donors, as compared to another CMV test.	≥ 99%
	Hospitalised patients ⁴	≥200	Potential limitations for specificity, if any, shall be identified
Cross-reactivity	Potentially cross-reacting ⁵ specimens	≥100 in total (e.g. RF+, related viruses or other infectious agents, pregnant women, etc.)	

¹ Including testing of other CMV parameters (e.g. CMV-IgM, avidity, immunoblot), or previous / follow-up samples for true sample status.

² Supplementary testing to confirm recent CMV infection (primary or re-infection): e.g. CMV-IgM, IgG-avidity, immunoblot analysis.

³ Corresponding to an initial number of 1000 donors at an assumed CMV prevalence of 60 %.

⁴ Including pre-transplant recipients.

⁵ Including related β -herpes viruses (HHV-6, HHV-7).

Table 2. Qualitative and quantitative NAT devices for CMV DNA

1. For target sequence amplification devices, a functionality control for each specimen (internal control) shall reflect the state of the art. This control shall as far as possible be used throughout the whole process, i.e. extraction, amplification/hybridisation, detection.
2. Genotype and/or subtype detection shall be demonstrated by appropriate primer or probe design validation and shall also be validated by testing characterised genotyped samples.
3. Potential cross-reactivity of non-target nucleic acid sequences shall be analysed by appropriate primer or probe design validation and shall also be validated by testing selected samples.
4. Results of quantitative NAT devices shall be traceable to international standards or calibrated reference materials, if available, and be expressed in international units utilised in the specific field of application.

Performance characteristic	Specimens	Specimen numbers, features, use	Acceptance criteria
Analytical sensitivity	WHO 1 st International Standard Human CMV DNA (09/162; 5,000,000 IU/vial) (or calibrated reference materials)	NAT sensitivity and NAT LOD shall be validated by dilution series of reference materials, testing of replicates (minimal 24) at different analyte concentrations, including those with transition from positive to negative results in the respective NAT assay. LOD shall be expressed as 95% positive cut-off value (IU/ml) after statistical analysis (e.g. Probit). ¹ quantitative NAT: definition of lower, upper quantification limit, precision, accuracy, 'linear' measuring range, 'dynamic range'. Reproducibility at different concentration levels	According to the state of the art

Diagnostic sensitivity CMV Strain sensitivity	Patient samples determined as CMV DNA positive by comparator device Dilution series of CMV positive cell cultures may serve as potential substitutes	Qualitative NAT: ≥ 100 Quantitative NAT: ≥ 100 dilution series for demonstration of quantification efficiencies	According to the state of the art
Diagnostic specificity	Blood donor specimens	Qualitative NAT: ≥ 500 Quantitative NAT: ≥ 100	According to the state of the art
Cross-reactivity	Potentially cross-reacting specimens	≥ 20 Including human samples positive for related human herpesviruses, e.g. EBV, HHV6, VZV Herpesvirus positive cell cultures may serve as potential substitutes	According to the state of the art
Carry-over	High CMV DNA positive; CMV DNA negative	At least five runs with alternating high-positive and negative specimens shall be performed during robustness studies. The virus titres of the high positive samples shall be representative of high virus titres occurring naturally.	According to the state of the art
Whole system failure rate	CMV DNA low positive	≥ 100 CMV DNA low-positive specimens shall be tested. These specimens shall contain a virus concentration equivalent to three times the 95 % positive cut-off virus concentration.	$\geq 99\%$ positive

¹ Reference: European Pharmacopeia 9.0, 2.6.21 Nucleic acid amplification techniques, Validation.

ANNEX X

COMMON SPECIFICATIONS FOR DEVICES INTENDED FOR DETECTION OR QUANTIFICATION OF MARKERS OF EPSTEIN-BARR VIRUS (EBV) INFECTION

Scope

This Annex applies to devices intended for detection or quantification of markers of Epstein-Barr virus (EBV) infection.

Table 1 applies to first-line assays for IgG antibodies against viral capsid antigen of EBV (anti-EBV VCA IgG).

Table 2 applies to qualitative and quantitative NAT devices for EBV DNA.

Table 1: First-line assays: anti-EBV VCA IgG

Performance characteristic	Specimens	Anti-EBV VCA IgG	Acceptance criteria
Diagnostic sensitivity	Positive specimens	≥400 including specimens from recent and past EBV infection, low and high positive titre samples	≥ 99% for confirmable past infection ¹ ; overall sensitivity including recent infection ² shall be equivalent to other CE-marked tests
	Seroconversion panels	To be tested when available	diagnostic sensitivity during seroconversion shall represent the state of the art
Analytical sensitivity	Standards	International reference reagents, when available	
Diagnostic specificity	Negative specimens	≥ 200 ³ EBV negatives from unselected donors as compared to another EBV test.	≥ 99%
	Hospitalised patients ⁴	≥200	Potential limitations for specificity, if any, shall be identified
Cross-reactivity	Potentially cross-reacting specimens	≥100 in total (e.g. RF+, related viruses or other infectious agents, pregnant women, etc.)	

¹ Including other EBV markers and parameters (e.g. VCA-IgM, EBNA-1 IgG, immunoblot) to assess the true specimen status.

² Supplementary testing to confirm recent EBV infection: e.g. VCA-IgM, IgG-avidity, immunoblot analysis.

³ At an assumed EBV prevalence of 80 % corresponding to an initial number of 1000 donors.

⁴ Including pre-transplant recipients.

Table 2. Qualitative and quantitative NAT devices for EBV DNA

1. For target sequence amplification devices, a functionality control for each specimen (internal control) shall reflect the state of the art. This control shall as far as possible be used throughout the whole process, i.e. extraction, amplification/hybridisation, detection.
2. Genotype and/or subtype detection shall be demonstrated by appropriate primer or probe design validation and shall also be validated by testing characterised genotyped samples.
3. Potential cross-reactivity of non-target nucleic acid sequences shall be analysed by appropriate primer or probe design validation and shall also be validated by testing selected samples.
4. Results of quantitative NAT devices shall be traceable to international standards or calibrated reference materials, if available, and be expressed in international units utilised in the specific field of application.

Performance characteristic	Specimens	Specimen numbers, features, use	Acceptance criteria
Analytical sensitivity	WHO 1 st International Standard Human EBV DNA (09/260; 5,000,000 IU/vial) (or calibrated reference materials)	NAT sensitivity and NAT LOD shall be validated by dilution series of reference materials, testing of replicates (minimal 24) at different analyte concentrations, including those with transition from positive to negative results in the respective NAT assay. LOD shall be expressed as 95% positive cut-off value (IU/ml) after statistical analysis (e.g. Probit). ¹ quantitative NAT: definition of lower, upper quantification limit, precision, accuracy, 'linear' measuring range, 'dynamic range'. Reproducibility at different concentration levels	According to the state of the art

Diagnostic sensitivity EBV strain sensitivity	Patient samples determined as EBV DNA positive by comparator device Dilution series of EBV positive cell cultures may serve as potential substitutes	Qualitative NAT: ≥ 100 Quantitative NAT: ≥ 100 dilution series for demonstration of quantification efficiencies	
Diagnostic specificity	Negative specimens	Qualitative NAT: ≥ 500 Quantitative NAT: ≥ 100	According to the state of the art
Cross-reactivity	Potentially cross-reacting specimens	≥ 20 Including human samples positive for related human herpesviruses, e.g. CMV, HHV6, VZV Herpesvirus positive cell cultures may serve as potential substitutes	According to the state of the art
Carry-over	High EBV DNA positive; EBV DNA negative	At least five runs with alternating high-positive and negative specimens shall be performed during robustness studies. The virus titres of the high positive samples shall be representative of high virus titres occurring naturally.	According to the state of the art
Whole system failure rate	EBV DNA low positive	≥ 100 EBV DNA low-positive specimens shall be tested. These specimens shall contain a virus concentration equivalent to three times the 95 % positive cut-off virus concentration.	$\geq 99\%$ positive

¹ Reference: European Pharmacopeia 9.0, 2.6.21 Nucleic acid amplification techniques, Validation.

ANNEX XI

COMMON SPECIFICATIONS FOR DEVICES INTENDED FOR DETECTION OF MARKERS OF *TREPONEMA PALLIDUM* INFECTION

Scope

This Annex applies to devices intended for detection of *Treponema pallidum* (*T. pallidum*).

Table 1 applies to first-line assays for antibodies against *T. pallidum* (anti-*T.pallidum*).

Table 2 applies to confirmatory and supplemental anti-*T.pallidum* assays.

Table 1. First-line assays: anti-*T.pallidum*

Performance characteristic	Specimens	Specimen number, features, use	Acceptance criteria
Diagnostic sensitivity	Positive specimens	≥200 positive samples in total, at different stages of the infection if available, including high positive and weakly positive samples, identified as positive by at least two different serological tests (one of which is an enzyme immunoassay) for different antibodies to <i>T.pallidum</i>	≥99.5% overall sensitivity
	Seroconversion panels	At least 1 seroconversion panel, ≥1 if possible, including individual samples from the early infection phase	Diagnostic sensitivity during seroconversion shall represent the state of the art.
Analytical sensitivity	Standards	WHO international standards NIBSC code 05/132, when available	
Diagnostic specificity	Unselected blood donors (including first-time donors) ¹	≥5000	≥99,5%
	Hospitalised patients	≥200	Potential limitations for specificity, if any, shall be identified
Cross-reactivity	Potentially cross-reacting blood- specimens	≥100 in total including the following specimens: positive for <i>Borrelia burgdorferi sensu lato</i> confirmed by IgG immunoblot; anti-HIV positive; RF+; other related microbial/infectious agents; systemic lupus erythematosus (SLE) patients; antiphospholipid antibody positive; pregnant women etc.	

¹ Blood donor populations shall be investigated from at least two blood donation centres and consist of consecutive blood donations, which have not been selected to exclude first-time donors.

Table 2. Confirmatory and supplemental assays: anti-*T.pallidum*

Performance characteristic	Specimens	Specimen number, features, use	Acceptance criteria
Diagnostic sensitivity	Positive specimens	≥300 positive samples at different stages of the infection (early primary syphilis, secondary stage, and during late syphilis) including high positive samples, 50 weakly positive samples, by at least two different serological tests (one of which is an enzyme immunoassay) for different antibodies to <i>T.pallidum</i>	99% identification as “confirmed positive” or “indeterminate”
	Seroconversion panels	At least 1 seroconversion panel , ≥1 if possible, including individual samples from the early infection phase	diagnostic sensitivity during seroconversion shall represent the state of the art
Analytical sensitivity	Standards	WHO international standards NIBSC code 05/132	
Diagnostic specificity	Blood donors	≥200	≥ 99%;
	Clinical samples	≥200	Potential limitations for specificity, if any, shall be identified
Cross-reactivity	Potentially cross-reacting specimens	≥50 in total, including samples from pregnant women and samples with indeterminate results in other confirmatory assays.	

ANNEX XII

COMMON SPECIFICATIONS FOR DEVICES INTENDED FOR DETECTION OR QUANTIFICATION OF MARKERS OF *TRYPANOSOMA CRUZI* INFECTION

Scope

This Annex applies for devices intended for detection or quantification of markers of *Trypanosoma cruzi* (*T. cruzi*) infection.

Table 1 applies to first-line assays for antibodies against *T. cruzi* (anti-*T. cruzi*).

Table 2 applies to confirmatory and supplemental anti-*T. cruzi* assays.

Table 3 applies to qualitative and quantitative NAT devices for *T. cruzi* DNA.

Table 1. First-line assays: anti-*T. cruzi*

Performance characteristic	Specimens	Specimen number, features, use	Acceptance criteria
Diagnostic sensitivity	Positive specimens	<p>≥400 positive samples, including highly positive confirmed by at least two different serological tests for different antibodies to <i>T. cruzi</i>.</p> <p>Of those 400, ≥25 parasite positive samples, which have been confirmed by direct detection.</p>	99.5% overall sensitivity
	Seroconversion panels	To be defined when available	Diagnostic sensitivity during seroconversion shall represent the state of the art
Analytical sensitivity	Standards	<p>WHO international standards</p> <p>NIBSC code: 09/186</p> <p>NIBSC code: 09/188</p>	
Diagnostic specificity	Unselected donors (including first-time donors) ¹	≥5000	≥99,5%
	Hospitalised patients	≥200	Potential limitations for specificity, if any, shall be identified

Cross-reactivity	Potentially cross-reacting specimens	≥100 in total including the following specimens: positive for anti- <i>Toxoplasma gondii</i> ; at least 5 specimens positive for anti- <i>Leishmania</i> ; RF+; related microbial agents or other infectious agents; SLE patients; antiphospholipid antibody positive patients; pregnant women, etc.	
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¹ Blood donor populations shall be investigated from at least two blood donation centres and consist of consecutive blood donations, which have not been selected to exclude first-time donors.

Table 2. Confirmatory and supplemental assays: anti-*T. cruzi*

Performance characteristic	Specimens	Specimen number, features, use	Acceptance criteria
Diagnostic sensitivity	Positive specimens	≥300 positive specimens, including highly positive confirmed by at least two different serological tests for different antibodies to <i>T. cruzi</i> . Of those 300, ≥25 parasite positive samples, which have been confirmed by direct detection.	≥99% identification as “confirmed positive” or “indeterminate”
	Seroconversion panels	As available	Diagnostic sensitivity during seroconversion shall represent the state of the art, if applicable
Analytical sensitivity	Standards	WHO international standards NIBSC code: 09/186 NIBSC code: 09/188	
Diagnostic specificity	Negative specimens	≥200	≥99%
	Clinical samples	≥200	Potential limitations for specificity, if any, shall be identified
Cross-reactivity	Potentially cross-reacting specimens	≥50, including samples from pregnant women and samples with indeterminate results in other confirmatory assays	

Table 3: NAT devices for *T. cruzi* DNA

1. For target sequence amplification devices, a functionality control for each specimen (internal control) shall reflect the state of the art. This control shall as far as possible be used throughout the whole process, i.e. extraction, amplification/hybridisation, detection.
2. Genotype and/or subtype detection shall be demonstrated by appropriate primer or probe design validation and shall also be validated by testing characterised genotyped samples.
3. Potential cross-reactivity of non-target nucleic acid sequences shall be analysed by appropriate primer or probe design validation and shall also be validated by testing selected samples.
4. Results of quantitative NAT devices shall be traceable to international standards or calibrated reference materials, if available, and be expressed in international units utilised in the specific field of application.

Performance characteristic	Specimens	Specimen number, features, use	Acceptance criteria
Analytical sensitivity	Characterized in-house reference preparation (as long as international reference materials are not available)	NAT sensitivity and NAT LOD shall be validated by dilution series of reference materials, testing of replicates (minimal 24) at different analyte concentrations, including those with transition from positive to negative results in the respective NAT assay. LOD shall be expressed as 95% positive cut-off value (IU/ml) after statistical analysis (e.g. Probit). ¹	According to the state of the art
Diagnostic sensitivity: different <i>T. cruzi</i> strains / isolates	Patient samples from different regions determined as <i>T. cruzi</i> DNA positive by comparator device; sequence variants	≥100 Dilution series of <i>T. cruzi</i> positive cell cultures (isolates) or <i>T. cruzi</i> positive materials from animal models may serve as potential substitutes	According to the state of the art
Diagnostic specificity	Negative specimens	≥100	According to the state of the art

¹ Reference: European Pharmacopeia 9.0, 2.6.21 Nucleic acid amplification techniques, Validation.

Cross-reactivity	Potentially cross-reacting specimens	≥10 human samples positive for other parasites, e.g. <i>Plasmodium</i> species, <i>Trypanosoma brucei</i> . Positive cell cultures may serve as potential substitutes	According to the state of the art
Carry-over		At least five runs with alternating high-positive and negative specimens shall be performed during robustness studies. The <i>T.cruzi</i> titres of the high positive samples shall be representative of high <i>T.cruzi</i> titres occurring naturally.	According to the state of the art
Whole system failure rate		≥100 <i>T.cruzi</i> DNA low-positive specimens shall be tested. These specimens shall contain a <i>T.cruzi</i> concentration equivalent to three times the 95 % positive cut-off <i>T.cruzi</i> concentration.	≥99% positive

ANNEX XIII

COMMON SPECIFICATIONS FOR DEVICES INTENDED FOR DETECTION OR QUANTIFICATION OF MARKERS OF SEVERE ACUTE RESPIRATORY SYNDROME CORONAVIRUS 2 INFECTION

Scope

This Annex applies to devices intended for detection or quantification of markers of severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) infection.

Table 1 applies to the following first-line assays (including rapid tests) for antibodies against SARS-CoV-2 (anti-SARS-CoV-2): total antibody, IgG-only, IgG combined with IgM and/or IgA.

Table 2 applies to first-line assays (including rapid tests) for detection of anti-SARS-CoV-2 IgM and/or IgA.

Table 3 applies to confirmatory or supplemental assays for anti-SARS-CoV-2.

Table 4 applies to antigen SARS-CoV-2 tests, including rapid antigen tests.

Table 5 applies to NAT assays for SARS-CoV-2 RNA.

Table 6 applies to SARS-CoV-2 antigen self-tests which have already undergone a performance evaluation for professional use.

Table 7 applies to SARS-CoV-2 antibody self-tests which have already undergone a performance evaluation for professional use.

Table 1: First-line assays (including rapid tests) for anti-SARS-CoV-2: total antibody, IgG-only, IgG combined¹ with IgM and/or IgA

Performance characteristic	Specimen	Anti-SARS-CoV-2 total antibody, IgG, IgG combined	Acceptance criteria
Diagnostic sensitivity	Positive specimens	≥ 400 including samples from early infection and post seroconversion ² (within the first 21 days and after 21 days following the onset of symptoms); including samples from asymptomatic or subclinical and mildly symptomatic (outpatient treatment) individuals; including samples with low and high titers; including samples from vaccinated individuals where appropriate ³ ; consideration of genetic variants	$\geq 90\%$ sensitivity ⁴ for samples taken >21 days after onset of symptoms ⁵ ; overall sensitivity including the early infection phase shall be comparable to other CE marked ⁶ tests
	Seroconversion panels	As far as available	Seroconversion sensitivity comparable to other CE-marked tests
Analytical sensitivity	Reference preparations	WHO International Standard (IS) for anti- SARS-CoV-2 (NIBSC code 20/136);	IS: for titre determinations / quantitative ⁷ result output;

		WHO International Reference Panel (RP) for anti-SARS-CoV-2 antibodies (NIBSC codes 20/140, 20/142, 20/144, 20/148, 20/150)	RP: all antibody assays
Specificity	Negative specimens ⁸	≥400 samples from non-infected and non-vaccinated individuals ⁹	>99% specificity ¹⁰
		≥200 hospitalised patients (without SARS-CoV-2 infection)	Potential limitations for specificity shall be determined
		≥100 in total potentially interfering (e.g. rheumatoid factor, pregnant women, etc.) and cross-reacting blood specimens: including antibodies against endemic human coronaviruses 229E, OC43, NL63, HKU1 and other pathogens of respiratory diseases such as influenza A, B, RSV etc.	

¹ Performance claim of the combined overall result; for devices with separate claims for IgM and/or IgA, see table 2.

² Details on the time interval between sampling and onset of symptoms (or time of infection, if available) shall be provided.

³ The manufacturer shall provide a justification of the suitability and timing for sensitivity evaluation of the relevant antibodies in vaccinated individuals.

⁴ Based on confirmed positive SARS-CoV-2 NAT result.

⁵ Claims for sensitivity shall be specified in relation to the time between sampling after symptom onset or the initial PCR diagnosis and the test.

⁶ CE marked under Regulation (EU) 2017/746 as class D, if available.

⁷ This applies to quantitative assays if they are also first-line assays.

⁸ Negative specimens shall be from individuals with no history of SARS-CoV-2 infection (if available pre-pandemic).

⁹ Individuals vaccinated with an antigen different from that used in the device may be included, if appropriate.

¹⁰ False positive results shall be resolved by re-testing in other SARS-CoV-2 serologic assays, if necessary with different test design and antigen coating compared to the initial test, and/or confirmatory testing.

Table 2: First-line assays (including rapid tests) for anti-SARS-CoV-2: IgM and/or IgA detection

Performance characteristic	Specimen	Anti-SARS-CoV-2 IgM and/or IgA	Acceptance criteria
Diagnostic sensitivity	Positive specimens	≥200 ¹ samples ² with a significant proportion from the	≥80% sensitivity ⁴ for samples taken during the first 21 days after symptom onset ⁵ ;

		early phase of the infection (within 21 days after onset of symptoms) compared to samples past seroconversion (>21 days after onset of symptoms); including samples from asymptomatic, subclinical, mildly symptomatic (outpatient treatment) individuals; including freshly ³ vaccinated individuals if appropriate; consideration of genetic variants	overall sensitivity shall be comparable to other CE marked ⁶ tests of the same type (i.e. IgM and/or IgA)
	Seroconversion panels	As far as available	Seroconversion sensitivity comparable to other CE-marked tests
Analytical sensitivity	Standards	N/A	N/A
Specificity	Negative specimens ⁷	≥200 samples from non-infected and non-vaccinated individuals ⁸	≥98% specificity ⁹
		≥100 from hospitalised patients (without SARS-CoV-2 infection)	Potential limitations for specificity shall be determined
		≥100 in total potentially interfering (e.g. rheumatoid factor, pregnant women, etc.) and cross-reacting blood specimens; antibodies against endemic human coronaviruses 229E, OC43, NL63, HKU1 and other pathogens of respiratory diseases such as influenza A, B, RSV etc.	

¹ In case of devices detecting both IgM and IgA, 200 per marker IgM and IgA.

² Details on the time interval between sampling and onset of symptoms (or time of infection, if available) shall be provided.

³ The manufacturer shall provide a justification of the suitability and timing for sensitivity evaluation of IgM and IgA in vaccinated individuals.

⁴ Based on confirmed positive SARS-CoV-2 NAT result.

⁵ Claims for sensitivity shall be specified in relation to the time between sampling after symptom onset or the initial PCR diagnosis and the test.

⁶ CE marked under Regulation (EU) 2017/746 as class D, if available.

⁷ Negative specimens shall be from individuals with no history of SARS-CoV-2 infection (if available pre-pandemic).

⁸ Individuals vaccinated with an antigen different from that used in the device may be included, if appropriate.

⁹ False positive results shall be resolved by re-testing in other SARS-CoV-2 serologic assays, if necessary with different test design and antigen coating compared to the initial test, and/or confirmatory testing. Clarification of false positive results may additionally include testing for presence of other anti-SARS-CoV-2 antibody types (IgA, IgG, total antibody).

Table 3: Confirmatory or supplemental¹ assays for anti-SARS-CoV-2

Performance characteristic	Specimen	Anti-SARS-CoV-2	Acceptance criteria
Diagnostic sensitivity	Positive specimens	≥200 including samples pre and post seroconversion (within the first 21 days and after 21 days following the onset of symptoms)	Correct determination as “positive” (or “indeterminate”)
	Seroconversion panels/ low titre panels	as far as available	
Analytical sensitivity	Standards	N/A	N/A
Diagnostic specificity	Negative specimens ²	≥200 from non-infected / non-vaccinated population	No false positive results; correct determination as “negative” (or “indeterminate”)
		≥200 from hospitalised patients (without SARS-CoV-2 infection) ≥50 potentially interfering and cross-reacting samples in total: antibodies against endemic human coronaviruses 229E, OC43, NL63, HKU1 and other pathogens of respiratory diseases such as influenza A, B, RSV etc.; including samples with indeterminate or false positive results in other anti-SARS-CoV-2 assays	

¹ E.g. immunoblot providing antigens different from those used in the initial antibody test.

² Negative specimens shall be from individuals with no history of SARS-CoV-2 infection (if available pre-pandemic).

Table 4: Antigen assays (including rapid tests): SARS-CoV-2

Performance characteristic	Specimen	SARS-CoV-2 antigen	Acceptance criteria
Diagnostic sensitivity	Positive specimens	$\geq 100^1$ NAT positive samples ² from early infection within the first 7 days after symptom onset ³ ; samples shall represent naturally occurring viral loads ⁴ ; consideration of genetic variants ⁵ ; consideration of variations in specimen collection and/or specimen handling ⁶	Detection of >80% (rapid tests); detection of >85% (lab-based assays ⁷); relative to SARS-CoV-2-NAT ^{8,9}
Analytical sensitivity	Standards	As soon as available	Establishment of a LOD ¹⁰
Diagnostic specificity	Negative specimens	≥ 300 from non-infected individuals	Specificity >98% (rapid tests) Specificity >99% (lab-based assays ⁷)
		≥ 100 from hospitalised patients ≥ 50 potentially interfering and cross-reactive samples in total: including virus-positive samples of endemic human coronaviruses 229E, OC43, NL63, HKU1; influenza A, B, RSV, and other pathogens of respiratory diseases, eligible for differential diagnosis; including bacteria ¹¹ present in the sampling area	Potential limitations for specificity shall be determined

¹ If the device is intended to be used for more than one specimen type, 100 samples shall be required for each specimen type. If this is not possible in exceptional circumstances (e.g. if specimen collection is very invasive), the manufacturer shall provide a justification and evidence of matrix equivalence.

² Sampling shall be matched for antigen and NAT testing, e.g., two simultaneous samples from each individual or optimally NAT- and antigen testing from the same sample (e.g. from the eluate of one swab); the buffer/transport medium shall be compatible for both NAT and antigen testing; any volume change in the buffer/medium for sample uptake different from that of the proprietary assay, and/or between antigen and NAT test shall be clearly communicated.

³ Or time of infection, if known, taking into account the incubation time.

⁴ I.e., without preselection; the viral loads and their distribution shall be shown, e.g. characterized by Ct-values of RT-PCR; or transformed into viral load per ml or sample, if applicable.

⁵ Depending on the design of the device and nature of the genetic variant. For the purpose of evaluation, at least 3 samples shall be represented for each genetic variant.

⁶ Specimen collection and extraction items such as swabs, extraction buffers, etc., shall be part of the evaluation. If proprietary sampling/sample preparation is not included in the test kit, test performance shall be investigated for an applicable range of sampling devices. If the sample is not tested immediately, e.g. after a certain transport time, stability of the antigen shall be investigated.

⁷ Other than rapid tests, i.e. formal laboratory-based assays e.g. enzyme immunoassay, automated tests, etc.

⁸ The sensitivity of $\geq 80\%$, $\geq 85\%$ respectively, shall be for all specimen types claimed. All claimed specimen types shall be compared with paired NAT results from nasopharyngeal specimens.

⁹ The relationship between antigen test performance and NAT shall be demonstrated; sensitivity may be shown relating to different viral load ranges and to the threshold of infectivity. The NAT and extraction method used shall be described.

¹⁰ Unless there is an available international standard, analytical sensitivity may be tested by dilution series of in-house virus preparations, comparatively with other antigen tests and NAT; if inactivated virus is used, the effect of inactivation and freeze/thawing on the antigen shall be investigated.

¹¹ E.g. staphylococci and streptococci expressing protein A or G.

Table 5: NAT assays for SARS-CoV-2 RNA

Performance characteristic	Specimen	SARS-CoV-2 RNA qualitative	SARS-CoV-2 RNA quantitative
Sensitivity			
Analytical sensitivity: LOD	WHO 1st International Standard SARS-CoV-2 RNA (NIBSC code 20/146; 7.70 Log ₁₀ IU/mL) Secondary standards calibrated against WHO IS	According to Ph. Eur. NAT validation guideline: several dilution series into borderline concentration; statistical analysis (e.g. Probit analysis) on the basis of at least 24 replicates; calculation of 95 % cut-off value	According to Ph. Eur. NAT validation guideline: several dilution series of calibrated reference preparations into borderline concentration; statistical analysis (e.g. Probit analysis) on the basis of at least 24 replicates; calculation of 95 % cut-off value as LOD
Quantification limit; quantification features	WHO 1st International Standard SARS-CoV-2 RNA (NIBSC code 20/146; 7.70 Log ₁₀ IU/mL) Secondary standards calibrated against WHO IS		Dilutions (half-log ₁₀ or less) of calibrated reference preparations; determination of lower, upper quantification limit, LOD, precision, accuracy, “linear” measuring range, “dynamic range”. Synthetic target may be used as secondary standard to achieve

			higher concentration levels. Reproducibility at different concentration levels to be shown
Diagnostic sensitivity: different SARS-CoV-2 RNA strains	Patient samples determined as SARS-CoV-2 RNA positive by comparator device from different regions and outbreak clusters; sequence variants Dilution series of SARS-CoV-2 positive cell cultures (isolates) may serve as potential substitutes	$\geq 100^1$	
Quantification efficiency	SARS-CoV-2 RNA positive patient samples from different regions and outbreak clusters; sequence variants with quantitative values obtained by comparator device Dilution series of SARS-CoV-2 RNA positive cell cultures may serve as potential substitutes		≥ 100
Inclusivity	<i>In silico</i> analysis ² ; at least two independent target gene regions in one test run (dual-target design)	Evidence of suitable assay design: primer/probe sequence alignments with published SARS-CoV-2 sequences	Evidence of suitable assay design: primer/probe sequence alignments with published SARS-CoV-2 sequences
Specificity			
Diagnostic specificity	SARS-CoV-2 RNA negative human specimens	≥ 500	≥ 100
<i>In silico</i> analysis ²		Evidence of suitable assay design evidence (sequence alignments); regular check of primer/probe sequences against sequence data bank entries	Evidence of suitable assay design evidence (sequence alignments); regular check of primer/probe sequences against sequence data bank entries
Potential cross reaction	samples positive (various concentrations) for related human coronaviruses 229E,	≥ 20 in total	≥ 20

	HKU1, OC43, NL63, MERS coronavirus; SARS CoV-1 if available; Influenza virus A, B; RSV; <i>Legionella pneumophila</i> ; positive cell cultures may serve as potential substitutes		
Robustness			
Cross contamination		At least 5 runs using alternating high positive and negative samples. The virus titres of the high positive samples shall be representative of high virus titres occurring naturally.	At least 5 runs using alternating high positive (known to occur naturally) and negative samples
Inhibition		Internal control preferably to go through the whole NAT procedure	Internal control preferably to go through the whole NAT procedure
Whole system failure rate leading to false negative results: 99/100 assays positive		≥100 samples virus-spiked with 3 × the 95 % positive cut-off concentration (3 x LOD)	≥100 samples virus-spiked with 3 × the 95 % positive cut-off concentration (3 x LOD)

¹If the device is intended to be used for more than one specimen type, 100 samples shall be required for each specimen type. If this is not possible in exceptional circumstances (e.g. if specimen collection is very invasive), the manufacturer shall provide a justification and evidence of matrix equivalence.

² The manufacturer shall define frequency and document evidence of regular surveillance checks against updated data bank entries in a post-market performance follow-up plan and report.

Table 6: Additional requirements for SARS-CoV-2 antigen self-tests¹

Performance characteristic	Specimens ²	Number of lay users	Criterion
Result interpretation	Interpretation of contrived tests ³ by lay users reflecting a range of results: <ul style="list-style-type: none"> • non-reactive • reactive • weak reactive⁴ • invalid 	≥100	Reading and interpretation of the contrived test results by 100 lay people; each lay person shall be subjected to read the specified range of result reactivity levels; determination of concordance of lay reading of the same tests by professional readers
Diagnostic sensitivity	Lay users that are known antigen positive ^{5,6}	≥30	In comparison to the true infectious status, i.e. by RT-PCR;

			concordance of results with the professional test
Diagnostic specificity	Lay users that do not know their status ⁵	≥60	Concordance of results with the professional test

¹ It is assumed that the underlying performance of the self-test has already been previously demonstrated with the evaluation/assessment of a professional test of the same design as the respective self-test under evaluation. In case for the self-use specimens in question there is no corresponding professional test variant, comparison shall be made with the standard specimen type (e.g. nasopharyngeal swabs for antigen test, serum or plasma for antibody test) of the corresponding professional test.

² For each self-use specimen type claimed with the device (e.g. nasal, sputum, saliva, whole blood, etc.).

³ Using whenever possible the original natural matrix of the respective specimen type.

⁴ A higher proportion of the samples shall be in the weak-positive range close to the cut-off or LOD of the test.

⁵ Individuals unaware of the professional diagnostic result prior to self-testing, and performing the entire test procedure from specimen collection and specimen pre-treatment (swab, buffer extraction, etc.) to reading.

⁶ Subjects up to about 7 days after symptom onset.

Table 7: Additional requirements for SARS-CoV-2 antibody self-tests¹

Performance characteristic	Specimens ²	Number of lay users	Criterion
Result interpretation	Interpretation of contrived tests ³ by lay users reflecting a range of results: <ul style="list-style-type: none"> • non-reactive • reactive • weak reactive⁴ • invalid 	≥100	Reading and interpretation of the contrived test results by 100 lay people; each lay person shall be subjected to read the specified range of result reactivity levels; determination of concordance of lay reading of the same tests by professional readers
Diagnostic sensitivity	Lay users that are known antibody positive ⁵	≥100	With previous history of initial PCR confirmed infection for SARS-CoV-2; in comparison to a previous confirmed antibody result; concordance of results with the professional test
Diagnostic specificity	Lay users that do not know their status ⁵	≥100	Concordance of results with the professional test

¹ It is assumed that the underlying performance of the self-test has already been previously demonstrated with the evaluation/assessment of a professional test of the same design as the respective self-test under evaluation. In case for the self-use specimens in question there is no corresponding professional test variant, comparison shall be made with the standard specimen type (e.g. nasopharyngeal swabs for antigen test, serum or plasma for antibody test) of the corresponding professional test.

² For each self-use specimen type claimed with the device (e.g. nasal, sputum, saliva, whole blood, etc.).

³ Using whenever possible the original natural matrix of the respective specimen type.

⁴ A higher proportion of the samples shall be in the weak-positive range close to the cut-off or LOD of the test.

⁵ Individuals unaware of the professional diagnostic result prior to self-testing, and performing the entire test procedure from specimen collection and specimen pre-treatment (swab, buffer extraction, etc.) to reading.