



Technical File - qPCR

FRIZ Biochem GmbH

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1 Meta Information

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1.1 Purpose

This Summary of Safety and Performance (SSP) is intended to provide public access to an up-to-date summary of the main aspects of the safety and performance of the IVD medical device GastroBac *direct* PCR.

The SSP is not intended to replace the Instructions For Use (IFU) as the main document to ensure the safe use of the device, nor is it intended to provide diagnostic or therapeutic suggestions to intended users.

The information in the next section is intended for professional users.

2 Summary of Safety and Performance (SSP) for professional users

2.1 Device identification and general information

Parameter	Value
Device trade name(s)	GastroBac <i>direct</i> PCR
Device reference number	FBC115 FBC115-Ax FBC115-Bx
Manufacturer´s name and address	FRIZ Biochem GmbH Floriansbogen 2-4 82061 Neuried Germany
Manufacturer´s single registration number (SRN)	DE-MF-000017533
Basic UDI-DI	426075389fbc115U5
European Medical Device Nomenclature (EMDN)	Device group: EMDN W0105: Infectious diseases Device type: EMDN W0105070203: Gram+ - Gram- - Fungi - Multiplex Assays
Risk class of the device	Class C
Indication whether it is a device for near-patient testing and/or a companion diagnostic	<input type="checkbox"/> Device for near-patient testing <input type="checkbox"/> Companion Diagnostic (CDx) <input checked="" type="checkbox"/> n.a.
Year when the first certificate was issued under Regulation (EU) 2017/746 covering the device	the device has not been certified under IVDR yet
Authorized representative, if applicable; name and the SRN	n.a.
NB´s name (the NB that will validate the SSP) and NB´s single identification number	Name of NB that will validate the SSP: TÜV SÜD Product Service GmbH NB´s single identification number: NB 0123

SSP reference number	FBC115GB818020379
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2.2 Intended use of the device

2.2.1 Intended purpose

DI Intended Purpose GastroBac

The GastroBac *direct* PCR is an assay for *in vitro* examination of bacterial DNA in stool samples to provide information to aid to diagnose patients under suspicion of gastrointestinal infection with *Campylobacter* spp., *Salmonella* spp., *Shigella* spp./enteroinvasive *Escherichia coli* (EIEC), *Yersinia enterocolitica* and/or *Yersinia pseudotuberculosis*.

The IVD medical device detects the DNA of the aforementioned pathogens by qualitative measurements based on qPCR and is intended for use in medical laboratories or health institutions by laboratory personnel specifically trained in qPCR and *in vitro* diagnostic techniques. It has to be used in combination with conventional nucleic acid extraction systems for DNA extraction and qPCR cyclers for detection and analysis.

2.2.2 Indication(s) and target population(s)

The GastroBac *direct* PCR may be applied for symptomatic patients under suspicion of gastrointestinal infection with *Campylobacter* spp., *Salmonella* spp., *Shigella* spp./Enteroinvasive *Escherichia coli* (EIEC), *Yersinia enterocolitica* and/or *Yersinia pseudotuberculosis*. Symptoms may include diarrhea, vomiting, abdominal pain, and fever. The target population encompasses subjects of all genders and ages. Gastrointestinal infection with the target organisms can affect individuals of any age, while children below 5 years of age are more frequently infected with *Campylobacter*, *Salmonella*, and *Yersinia*.

2.2.3 Limitation(s) and/or contra-indication(s)

There are no known contraindications or limitations to the target patient group. For detailed limitation on limitations and/or contra indications arising from associated interferences, cross-reactions, etc. see Instructions for Use available at www.frizbiochem.de.

2.3 Device description

2.3.1 Description of the device

DI Intended Purpose GastroBac

The GastroBac *direct* PCR is a multiplex qPCR test for the detection and differentiation of the following pathogens:

- *Campylobacter* spp. (*C. coli*, *C. jejuni*, *C. lari*, *C. upsaliensis*, and *C. hyointestinalis*)
- *Salmonella* spp. (*Salmonella enterica* subsp. *enterica* including *S. Typhimurium*, *S. Enteritidis*, *S. Typhi*, *S. Paratyphi A/B/C*)

- *Shigella* spp. (*S. flexneri*, *S. boydii*, *S. dysenteriae*, *S. sonnei*) and EIEC
- *Yersinia* (*Y. enterocolitica* and *Y. pseudotuberculosis*)

A separate Internal Control (IC) is added to each sample prior to DNA extraction and serves as control for nucleic acid isolation from the biological specimen as well as for amplification.

The nucleic acid eluate is added to the ready-to-use reaction solution that contains all reagents necessary for qPCR. The analysis can be performed on a qPCR cyclor.

The GastroBac *direct* PCR contains primer and probes specific for the aforementioned targets as well as the Internal Control. The probes are each labelled with fluorescent reporter dyes and a second dye that serves as a quencher and suppresses the fluorescence signals of intact probes.

The analysis is performed by determining Ct (cycle threshold) values. The Ct value describes the cycle in which the signal rises above a certain threshold for the first time. The more target copies are present in the sample, the lower the Ct value.

2.3.2 Description of the components

Each kit contains the following vials which are sufficient for 96 reactions (see *Table 1*).

Table 1: GastroBac direct PCR package content

Material	Lid colour	#vials; volume	#rxns.	Comment
Solution A	green	1x; 1050 µL	96	Reaction mix (buffer, enzymes, primer and probes)
Internal Control	blue	1x; 400 µL	96	Internal Control (artificial nucleic acid target)
Positive Control	red	1x; 50 µL	4	Plasmid DNA of target gene regions

2.3.3 Previous generation(s) or variants

There are no previous generations of the IVD medical device.

The GastroBac *direct* PCR is available in the following variants:

Table 2: GastroBac direct PCR configurations

Article number	Configuration	Channel of Internal Control
FBC115-Ax	suitable for use on a qPCR instrument with at least 5 detection channels (Cyan500/FAM/HEX/Red610/Cy5)	Cyan500
FBC115-Bx	suitable for for use on a qPCR instrument with at least 5 detection channels (FAM/HEX/Red610/Cy5/Cy5.5)	Cy5.5

2.3.4 Accessories

Additional components required that are not provided with the kit:

- qPCR cycler (with at least 5 detection channels)
- Disposable protective gloves, powder-free
- PCR reaction tubes/microtiter plate plus lids/adhesive optical film
- Pipettes
- sterile filter-tips for PCR testing (DNA/RNA-free)
- Table centrifuge
- DNA extraction kit (IVD-1049 chemagic™ Pathogen NA gDNA Kit H96 by PerkinElmer chemagen Technologie GmbH, MagNA Pure 96 DNA and Viral NA Small Volume Kit by Roche Life Science, or similar devices)
- Negative Control (no-template control, molecular grade water, or any other negative control according to the laboratory's standard procedure)

2.3.5 Other devices and products to be used in combination

The GastroBac *direct* PCR is intended to be used in combination with conventional nucleic acid extraction systems for DNA extraction and qPCR cyclers for detection and analysis.

2.4 List of harmonized standards and common specifications (CS) applied

Standard	Title
BS EN ISO 23640:2015	In vitro diagnostic medical devices. Evaluation of stability of in vitro diagnostic reagents
BS EN ISO 17511:2021	In vitro diagnostic medical devices. Requirements for establishing metrological traceability of values assigned to calibrators, trueness control materials and human samples
BS EN ISO 14971:2019+A11:2021	Medical devices. Application of risk management to medical devices
BS EN ISO 15223-1:2021	Medical devices. Symbols to be used with information to be supplied by the manufacturer – General requirements
BS EN ISO 13485:2016+A11:2021	Medical devices. Quality management systems. Requirements for regulatory purposes
BS EN ISO 20916:2024	In vitro diagnostic medical devices. Clinical performance studies using specimens from human subjects. Good study practice

2.5 Risks and warnings

2.5.1 Residual risks and undesirable effects

We successfully applied risk mitigation strategies. These measures reduced the probability category by at least one level without introducing new risks, thus we positively affected the benefit-risk ratio. After measure implementation all individual residual risks are accepted and consequently the overall residual risk is acceptable.

The GastroBac *direct* PCR fulfils the following parameters:

- compliance with the applicable general safety and performance requirements
- acceptable performance claims according to the state of the art and available diagnostic alternatives
- suitability of the device and its IFU for the intended users and use environment, including usability aspects

The residual risks are unlikely/moderate (probability/severity) or less.

Based on the clinical evidence and in alignment with the state of the art and the common specifications, GastroBac *direct* PCR achieves a clinical benefit by accurately detecting and differentiating DNA of *Campylobacter* spp., *Salmonella* spp., *Shigella* spp./EIEC, *Yersinia enterocolitica*, and *Yersinia pseudotuberculosis* in human clinical specimens (i.e., stool samples) and supporting diagnostic decision-making, taking into account additional clinical and laboratory information.

The clinical benefit is, to provide physicians with useful information with respect to patient management and therapy decisions, if applicable. This may help separate infected people from uninfected people and to isolate them from the rest of the population especially for high-risk patients.

The overall residual risk is in line with the risk policy and acceptable.

2.5.2 Warnings and precautions

The GastroBac *direct* PCR is intended for *in vitro* diagnostic use only. The test should only be performed by personnel trained in molecular diagnostic techniques. If the user makes substantial changes to the product or the application instructions, results may not correlate with the intended purpose.

- Before performing the test, read the entire instructions for use and follow them carefully. Deviations from the given test protocols can lead to invalid results.
- All patient samples must be treated as potentially infectious material.
- Discard sample and assay waste (that was in contact with patient material) according to your local, regional, or national safety regulations.
- The concentration specifications and incubation times of the manufacturers must be followed.
- Do not use the test beyond the expiration date.
- Do not use the test with opened or damaged packaging.
- Protect reagents from heat, moisture, and light.
- Do not replace or mix the reagents with reagents from other batches or other chemicals.
- Avoid contamination of the test by microorganisms and nucleases (DNases).
- Any carry-over of samples during handling and processing of the test may result in false positive test results.
- Good laboratory practice should be followed during the test.
- Use separated and segregated working areas for (1) sample preparation, (2) reaction setup and (3) amplification/detection activities. The workflow in the laboratory should proceed in unidirectional manner.
- Always wear disposable gloves in each area and change them before entering a different area.
- The test kits are intended for single use and must not be reused after performing qPCR reaction.
- If contamination of the qPCR cyclers is suspected, cleaning and maintenance must be carried out according to the system's manual.
- Safety Data Sheets (SDS) are available for download via <https://frizbiochem.de/downloads/>
- Pending EUDAMED entry, the Safety and Performance Summary (SSP) can be downloaded via <http://www.frizbiochem.de>

2.5.3 Other relevant aspects of safety

The chapter is not applicable as no field safety corrective actions (FSCA) and field safety notices (FSN) have been taken.

2.6 Summary of performance evaluation and post-market performance follow-up (PMPF)

2.6.1 Summary of scientific validity

To demonstrate the scientific validity of the GastroBac *direct* PCR, the following data sources were used:

- information (e.g., IFU) about the scientific validity of similar devices according to the State-of-the-Art report and
- scientific (peer-reviewed) literature

We selected PubMed (MEDLINE database) as the source for the literature search.

A separate literature search was performed for each analyte (*Campylobacter*, *Salmonella*, *Shigella*, *Yersinia*) considering the last 20 years (2004 – 2024). We defined the following exclusion criteria:

- The respective target organism was not considered.
- The respective target gene was not considered.
- The publication focuses on food safety.
- The publication investigates only samples from animals.
- The publication does not focus on PCR/NAT-based detection.
- The publication focuses on the prevalence/characterization of strains and/or antibiotic resistances.

The following table provides an overview of how many publications were excluded after abstract screening and how many publications were considered for demonstration of scientific validity.

Table 3: Results of the literature searches

Analyte	Literature search results	Excluded after abstract screening	Considered for scientific validity
<i>Campylobacter</i>	22	20	2
<i>Salmonella</i>	28	23	5
<i>Shigella</i>	24	12	12
<i>Yersinia</i>	28	16	12

In summary, it could be shown that qPCR is a powerful method for a fast and comprehensive diagnosis of patients with severe gastroenteritis. In general, rapid PCR assays show a sensitivity and specificity equivalent to that of culture.

Moreover, it could be shown that the chosen target genes underlying GastroBac *direct* PCR are appropriate targets for the detection of *Campylobacter* spp., *Salmonella* spp., *Shigella* spp./EIEC, *Yersinia enterocolitica*, and *Yersinia pseudotuberculosis*. All targets are well described in literature, and many of them are also used by similar products.

Finally, the scientific validity for the GastroBac *direct* PCR was demonstrated since the detection of the target genes of *Campylobacter* spp., *Salmonella* spp., *Shigella* spp./EIEC, *Yersinia enterocolitica*, and *Yersinia pseudotuberculosis* is verifiably associated with the infection of the respective pathogen.

2.6.2 Summary of performance data from equivalent device(s)

The chapter is not applicable as evidence of the underlying IVD medical device is not based on data from devices claimed to be equivalent to the device.

2.6.3 Summary of performance data from conducted studies prior to CE-marking

The chapter is not applicable as no performance studies have been conducted according to IVDR, Annex I, section 9 prior to CE-marking.

2.6.4 Summary of performance data from other sources

The chapter is not applicable as clinical evidence of the underlying IVD medical device is not based on data from other sources e.g., published experience gained by routine diagnostic testing.

2.6.5 Overall summary of the performance and safety

2.6.5.1 Analytical performance

GastroBac *direct* PCR test shows analytical specificity, analytical sensitivity, and precision that are similar to competitor devices. All analytical performance claims were able to be proven (*Table 4*).

Table 4: Overview of all applicable analytical performance parameters and the intended claims in comparison to the achieved results.

PE Analytical Performance Report GastroBac			
Analytical performance parameter	Claim	Result/s	Requirement fulfilled
Specimen collection, handling and stability	The GastroBac <i>direct</i> PCR is performed with nucleic acid that has been extracted from stool specimen. Quality and yield of DNA is highly dependent on the DNA extraction method used. The analytical performance is evaluated based solely on extracted DNA and thus independent of specimen collection, handling and stability.	N/A	<input checked="" type="checkbox"/> yes <input type="checkbox"/> no

Analytical sensitivity	see Limit of detection	<input checked="" type="checkbox"/> yes <input type="checkbox"/> no	
Analytical specificity	See Cross-reactivities, Endogenous and exogenous interference, Inclusivity, Competitive interference and Physical Interaction of primers and probes	<input checked="" type="checkbox"/> yes <input type="checkbox"/> no	
Cross-reactivities	<p>Cross-reactivity is tested with in silico analysis as well as with wet-lab testing. For the in silico analysis, the following acceptance criterion is applied:</p> <ul style="list-style-type: none"> • The homology between the cross-reactive microorganism and at least one primer sequence is < 80%. If both primer show a homology of $\geq 80\%$ the respective organism is also wet-lab tested. <p>For wet-lab testing the following acceptance criteria are applied:</p> <ul style="list-style-type: none"> • No false positive test results for tests with negative samples regarding the test analytes. • No false negative test results for tests with positive samples regarding the test analytes. 	<p>From the wet-lab tested potentially cross-reacting organisms, none gave a false positive nor false negative result. Therefore, cross-reactivity can be excluded.</p>	<input checked="" type="checkbox"/> yes <input type="checkbox"/> no

Endogenous and exogenous interferences	<p>Both positive and negative samples regarding the test analytes are tested with the spiked interfering substance. The following acceptance criteria are applied:</p> <ul style="list-style-type: none"> • No false positive test results for tests with negative samples regarding the test analytes. • No false negative test results for tests with positive samples regarding the test analytes. 	Tested substances at relevant concentrations did not interfere with the performance. There were no false positive nor false negative test results.	<input checked="" type="checkbox"/> yes <input type="checkbox"/> no
Inclusivity	<p>For inclusivity analysis, an in silico analysis is performed. The following acceptance criterion is applied:</p> <ul style="list-style-type: none"> • Frequency of sequences with exact match: minimum 95% of the target sequences <p>Otherwise, the respective isolate is wet-lab tested.</p>	The primer and probe sequences are suitable to detect the test analytes.	<input checked="" type="checkbox"/> yes <input type="checkbox"/> no
Competitive interference	<p>To analyse competitive interference, one test analyte is applied in a high concentration (e.g., 10^6 copies/mL), whereas the other test analytes are applied in a low concentration ($3 \times \text{LoD}$). The following acceptance criterion is applied:</p> <ul style="list-style-type: none"> • No false negative test results for all test analytes. 	<p>In most cases, a high concentration of spiking analyte up to 10^6 DNA copies/ml did not affect the detection of other analytes present at a low concentration ($3 \times \text{LoD}$). However, there was one exception. In the presence of 10^6 DNA copies/ml <i>Yersinia pseudotuberculosis</i>, one out of three <i>Salmonella enterica</i> samples was not detectable. In the presence of 10^5 DNA copies/ml <i>Yersinia pseudotuberculosis</i>, all <i>Salmonella enterica</i> replicates were detected.</p>	<input checked="" type="checkbox"/> yes <input type="checkbox"/> no (Note in the IFU about competitive interference)

Physical Interaction of primers and probes	The physical interaction of the different assay primers and probes are investigated with an in silico analysis. The following acceptance criteria are applied: <ul style="list-style-type: none"> • ΔG (kcal/mol) shall be > -9.0 and > -6.0 at 3' complementary base pairs. 	Potential dimers that do not fulfil the acceptance criteria were identified. Other parameters of the analytical performance studies show that potential dimers do not have an effect on the performance of the GastroBac <i>direct</i> PCR.	<input checked="" type="checkbox"/> yes <input type="checkbox"/> no
Accuracy	see trueness and precision		<input checked="" type="checkbox"/> yes <input type="checkbox"/> no
Trueness (bias)	No certified reference materials or reference measurement procedures are available. Thus, the comparison to the current clinical standard practice is addressed within the clinical performance evaluation.	N/A	<input checked="" type="checkbox"/> yes <input type="checkbox"/> no
Precision	see repeatability and reproducibility		<input checked="" type="checkbox"/> yes <input type="checkbox"/> no
Repeatability	The following acceptance criterion is applied: <ul style="list-style-type: none"> • CV $< 3\%$ 	In accordance with the acceptance criteria, the repeatability CV values ranged between 0.48% and 2.88%, depending on the analyte.	<input checked="" type="checkbox"/> yes <input type="checkbox"/> no
Reproducibility	The following acceptance criterion is applied: <ul style="list-style-type: none"> • CV $< 5\%$ 	In accordance with the acceptance criteria, the reproducibility CV values ranged between 0.98% and 3.61%, depending on the analyte.	<input checked="" type="checkbox"/> yes <input type="checkbox"/> no
Carryover/Cross-contamination	At least five runs using alternating high positive and negative specimens regarding the test analytes are performed. The following acceptance criteria are applied: <ul style="list-style-type: none"> • No false negative test results. • No false positive test results. 	In accordance with the acceptance criteria, there were no false negative nor false positive results.	<input checked="" type="checkbox"/> yes <input type="checkbox"/> no

Measuring range, Linearity	As the GastroBac <i>direct</i> PCR delivers a qualitative result (“detected” or “not detected”), the measuring range and linearity are not assessed.	N/A	N/A
Limit of detection	<p>The limit of detection is determined for each analyte separately. The following acceptance criterion is applied:</p> <ul style="list-style-type: none"> The limit of detection is \leq 10 copies/reaction for each analyte <p>To verify the limit of detection on a CFX Opus 96™ (Bio-Rad) cyclers, 20 replicates of each target at 1x LoD are run on a CFX Opus 96™. The following acceptance criterion is applied:</p> <ul style="list-style-type: none"> At least 19/20 replicates shall be positive for each target analyte (\geq 95% agreement) 	<p>The limit of detection in copies/reaction for the different analytes is:</p> <ul style="list-style-type: none"> Campylobacter: 0.383 (0.087 - 1.400) Salmonella: 1.195 (0.736 - 1.770) Shigella: 1.665 (0.407 - 5.058) EIEC: 4.160 (1.432 - 9.638) Yersinia enterocolitica: 3.546 (1.644 - 6.501) Yersinia pseudotuberculosis: 6.876 (2.523 - 15.488) <p>Moreover, at least 19/20 replicates were positive for each target analyte, when the LoD was verified on a CFX Opus 96™ (Bio-Rad) cyclers (\geq 95% agreement).</p>	<input checked="" type="checkbox"/> yes <input type="checkbox"/> no
Limit of quantitation	As the GastroBac <i>direct</i> PCR delivers a qualitative result (“detected” or “not detected”), the limit of quantitation is not.	N/A	N/A
Assay cut-off	As the GastroBac <i>direct</i> PCR delivers a qualitative result (“detected” or “not detected”), the assay cut-off is not assessed.	N/A	N/A
Metrological traceability of calibrator and control material values	The GastroBac <i>direct</i> PCR kit contains an internal control which is based on an artificial sequence. No suitable reference materials or reference measurement procedures of higher metrological order are available.	N/A	N/A

Stability	See Transport stability, Shelf life, and In-use stability		<input checked="" type="checkbox"/> yes <input type="checkbox"/> no
Transport stability	<p>Before and after the transport simulation, negative and positive samples regarding the test analytes are tested. The following acceptance criteria are applied:</p> <ul style="list-style-type: none"> • No false positive test results. • The agreement of the test results before and after the transport simulation is $\leq 10\%$ for each test analyte and the Internal Control. 	<p>In accordance with the acceptance criteria, Ct values deviated $\leq 10\%$ for each test analyte and the Internal Control after simulated transport.</p>	<input checked="" type="checkbox"/> yes <input type="checkbox"/> no
Shelf life	<p>The GastroBac <i>direct</i> PCR shall have a shelf life of 12 months when stored between temperatures from $-25\text{ }^{\circ}\text{C}$ to $-18\text{ }^{\circ}\text{C}$.</p> <p>Within the shelf-life study, positive and negative samples regarding the test analytes as well as the Positive Control are tested every month until 13 months. The following acceptance criteria are applied:</p> <ul style="list-style-type: none"> • No false positive test results. • The agreement (ΔCt) of the test results between t_0 and a measurement test point is $\leq 10\%$ for each test analyte and the Internal Control. 	<p>The shelf life study is still ongoing. A shelf life of 8 months for Solution A and IC could be confirmed.</p>	<input checked="" type="checkbox"/> yes <input type="checkbox"/> no

In-use stability	<p>The GastroBac <i>direct</i> PCR should be able to be thawed and refrozen 6 times. The following acceptance criteria are applied:</p> <ul style="list-style-type: none"> • No false positive test results • The agreement (ΔCt) of the test results between thawing cycle 1 and a later thawing cycle is $\leq 10\%$ for each test analyte and the Internal Control. <p>Moreover, the GastroBac <i>direct</i> PCR shall have an in-use stability up to 8 hours in a temperature range from 2°C to 8°C. The following acceptance criteria are applied:</p> <ul style="list-style-type: none"> • No false positive test results • The agreement (ΔCt) of the test results before and after storage at 4 °C is $\leq 10\%$ for each test analyte and the Internal Control. 	In accordance with the acceptance criteria, Ct values deviated $\leq 10\%$ for each test analyte and the Internal Control after 6 times thawing and refreezing as well as after 8 hours of storage at 4 °C.	<input checked="" type="checkbox"/> yes <input type="checkbox"/> no
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2.6.5.2 Clinical performance

For the evidence of clinical performance of the GastroBac *direct* PCR, we evaluated the parameters according to Annex I, section 9.1 b) of the IVDR. Diagnostic sensitivity and specificity data of the GastroBac *direct* PCR fulfill the acceptance criteria (Table 5). In conclusion, clinical performance data demonstrate clinical evidence of the GastroBac *direct* PCR.

Table 5: Overview of applicable clinical performance parameters and the intended claims in comparison to the achieved results.

PE Clinical Performance Report GastroBac

Clinical Performance Parameter	Acceptance criteria	Results
Diagnostic sensitivity	For each analyte, the diagnostic sensitivity is $\geq 90\%$	<p>For Campylobacter the diagnostic sensitivity is 100.0% (92.6 - 100.0)</p> <p>For Salmonella the diagnostic sensitivity is 100.0% (83.9 - 100.0)</p> <p>For Yersinia the diagnostic sensitivity is 100.0% (82.4 - 100.0)</p> <p>For Shigella the diagnostic sensitivity is 100.0% (83.2 - 100.0)</p>
Diagnostic specificity	For each analyte, the diagnostic specificity is $\geq 95\%$	<p>For Campylobacter the diagnostic specificity is 98.0% (89.4 - 100.0)</p> <p>For Salmonella the diagnostic specificity is 100.0% (92.9 - 100.0)</p> <p>For Yersinia the diagnostic specificity is 100.0% (92.9 - 100.0)</p> <p>For Shigella the diagnostic specificity is 100.0% (92.9 - 100.0)</p>

2.6.6 Ongoing or planned post-market performance follow-up (PMPF)

We use a Post-Market Performance Follow-Up (PMPF) Plan to define PMPF measures to be taken for the GastroBac *direct* PCR in order to keep the performance evaluation continually updated. The results of the PMPF activities are summarized and evaluated in the post-market performance follow-up report.

The following PMPF activities are planned:

- Research customer feedback, if applicable evaluate for trends and initiate action.
- Compile information from in-house quality control tests. If applicable, evaluate for trends and initiate action.
- Search for information on equivalent/comparative products and alternative procedures:
 - Review the scientific literature and other sources for relevant (clinical) performance and scientific data
 - Monitor relevant clinical guidelines from professional (medical) societies
 - Verify the performance data of equivalent and similar products, including searches in safety databases
- Search for changes in regulations, laws, initiate action if necessary
- Regular check of primer/probe sequences against sequence data bank entries
- Interlaboratory comparisons; further quality assurance programs

2.7 Metrological traceability of assigned values

The GastroBac *direct* PCR kit contains an internal control which is based on an artificial sequence. No suitable reference materials or reference measurement procedures of higher metrological order are available.

2.7.1 Explanation of the unit of measurement

The chapter is not applicable as no suitable reference materials or reference measurement procedures of higher metrological order are available.

2.7.2 Applied reference materials and/or reference measurement procedures

The chapter is not applicable as no suitable reference materials or reference measurement procedures of higher metrological order are available.

2.8 Suggested profile and training for users

We define the following user groups:

Table 6: User Groups

No.	Name of the User Group	Short Description
1	Healthcare Professional	tertiary/indirect user: collects and handles specimen, uses the results to derive an organizational and/or patient management decision
2	Laboratory Personnel	primary user: uses the GastroBac <i>direct</i> PCR for in vitro diagnostic analysis in a medical laboratory.
3	Laboratory Specialist	primary user: responsible for evaluation and release of the results

Table 7: User Group 1 “Healthcare Professional” as tertiary user

User group:	Healthcare Professional
Typical job title(s):	Physician, nurse, caregiver, pharmacist, pharmaceutical-technical assistant
Demographic characteristics (age, sex, language, specific physical attributes):	no specific gender typical age: between 20 and 67 years not necessarily a native speaker, but knows the required technical terminology
Expected qualification (education, degree, training):	medical or pharmaceutical education
Expected job experience (related to the product, similar products or IT in general):	No specific job experience related to the product required
Typical work environment (physical and organizational/social):	clinical environment (e.g. reception area), doctors' office

Core tasks:	1. Register a patient/specimen
	2. Collect a specimen
	3. Ship the collected specimen
	4. Interpret results to make an organizational and/or patient management decision
Typical equipment (used during performance of the tasks):	Swab and transport medium for specimen collection
Expected product training (with regard to the medical device)	no training required

Table 8: User Group 2 “Laboratory Personnel” as primary user

User group:	Laboratory Personnel
Typical job title(s):	medical/biological-laboratory assistant, laboratory technician
Demographic characteristics (age, sex, language, specific physical attributes):	no specific gender typical age: between 20 and 67 years not necessarily a native speaker, but knows the required technical terminology
Expected qualification (education, degree, training):	education as medical/biological-laboratory assistant or laboratory technician
Expected job experience (related to the product, similar products or IT in general):	Experience in molecular diagnostic techniques, i.e. nucleic acid extraction and qPCR
Typical work environment (physical and organizational/social):	medical laboratory
Core tasks:	1. Receive the specimen
	2. Register the patient/specimen
	3. Perform nucleic acid extraction according to the laboratory’s standard

	4. Perform the GastroBac <i>direct</i> PCR using a qPCR cyclers
Typical equipment (used during performance of the tasks):	qPCR cyclers disposable protective gloves PCR reaction tubes/microtiter plate plus lids/adhesive optical film Pipettes Pipette tips with filter (DNase/RNase-free) Table centrifuge DNA isolation kit
Expected product training (with regard to the medical device)	Training in qPCR and <i>in vitro</i> diagnostic techniques

Table 9: User Group 3 “Laboratory Specialist” as primary user

User group:	Laboratory Specialist
Typical job title(s):	specialist in laboratory medicine
Demographic characteristics (age, sex, language, specific physical attributes):	no specific gender typical age: between 30 and 67 years not necessarily a native speaker, but knows the required technical terminology
Expected qualification (education, degree, training):	university degree, specialist training for laboratory medicine
Expected job experience (related to the product, similar products or IT in general):	specialist training in infectious respiratory diseases
Typical work environment (physical and organizational/social):	medical laboratory
Core tasks:	Analyze and release the diagnostic report
Typical equipment (used during performance of the tasks):	qPCR cyclers with corresponding software LIMS software

Expected product training (with regard to the medical device)	Analysis and interpretation of the results
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3 Annex

3.1 Terms and Abbreviations

The following table lists document-specific abbreviations and definitions:

Term	Description	Reference
CDx	Companion diagnostic	n. a.
Companion diagnostic	Device which is essential for the safe and effective use of a corresponding medicinal product to: <ul style="list-style-type: none"> • identify, before and/or during treatment, patients who are most likely to benefit from the corresponding medicinal product; or • identify, before and/or during treatment, patients likely to be at increased risk of serious adverse reactions as a result of treatment with the corresponding medicinal product 	IVDR, Art. 2
Contraindication	Specific medical reasons for not using a particular IVD medical device in the usual way (e. g. a virus infection that interferes with the biomarker analysis, a urinary tract infection that affects the results of a urine test).	internal definition
DI	Device Identifier	n. a.
EMDN	European Medical Device Nomenclature	n.a.
EUDAMED	European Database on Medical Devices	n.a.
FSCA	Field Safety Corrective Action; Corrective action taken by a manufacturer for technical or medical reasons to prevent or reduce the risk of a serious incident in relation to a device made available on the market	IVDR, Art. 2
FSN	Field Safety Notice; Communication sent by a manufacturer to users or customers in relation to a field safety corrective action	IVDR, Art. 2

IFU	Instructions For Use; Information provided by the manufacturer to inform the user of a device's intended purpose and proper use and of any precautions to be taken	IVDR, Art. 2
Intended purpose	Use for which the IVD medical device is intended according to the data supplied by the manufacturer on the label, the IFU, or in promotional and sales materials and as specified by the performance evaluation.	IVDR, Art. 2
IVD	In Vitro Diagnostic	n. a.
IVDD	Directive 98/79/EC of the European Parliament and of the Council of 27 October 1998 on in vitro diagnostic medical devices	n. a.
IVDR	European Regulation 2017/476 on in vitro diagnostic medical devices	n. a.
Kit	Set of components that are packaged together and intended to be used to perform a specific in vitro diagnostic examination, or a part thereof	IVDR, Art. 2
Lay person	Individual who does not have formal education in a relevant field of healthcare or medical discipline	IVDR, Art. 2
Limitation	A situation/factor that (may) affect the results of an in-vitro diagnostic test (e.g. running a marathon before Ca ²⁺ -measurement due to untypically high Ca ²⁺ -level) or for which the test has not been clinically validated (e.g. usage for children, usage during pregnancy).	internal definition
MDSW	Medical device software	MDCG 2019-1 1
n. a.	not applicable	n. a.
NB	Notified Body; conformity assessment body designated in accordance with this Regulation	IVDR, Art. 2
Near-patient testing	Any device that is not intended for self-testing but is intended to perform testing outside a laboratory environment, generally near to, or at the side of, the patient by a health professional	IVDR, Art. 2
PEP	Performance evaluation plan	n. a.
PER	Performance evaluation report	n. a.

Performance of a device	Ability of a device to achieve its intended purpose as claimed by the manufacturer. It consists of the analytical and, where applicable, the clinical performance supporting that intended purpose	IVDR, Art. 2
PMPF	Post-Market Performance Follow-up	n. a.
Scientific validity	Association of an analyte with a clinical condition or a physiological state	IVDR, Art. 2
Self-testing	Any device intended by the manufacturer to be used by lay persons, including devices used for testing services offered to lay persons by means of information society services	IVDR, Art. 2
SRN	Single Registration Number	n. a.
SSP	Summary of Safety and Performance	n. a.
SVR	Scientific validity report	n. a.
TD	Technical documentation	n. a.
UDI	Unique Device Identification; Series of numeric or alphanumeric characters that is created through internationally accepted device identification and coding standards and that allows unambiguous identification of specific devices on the market	IVDR, Art. 2

3.2 Revision history

SSP revision number	Date issued	Change description	Revision validated by the Notified Body	
4	2025-08-04	Initial creation	<input type="checkbox"/>	Yes Validation language: ENG
5	2025-12-15	See Meta Information section	<input type="checkbox"/>	Yes Validation language: ENG
6	2026-01-07	See Meta Information	<input type="checkbox"/>	Yes Validation language: ENG