

Rx Only

cobas[®] Cdiff Test

for use on the cobas[®] 4800 System

For in vitro diagnostic use



cobas [®] 4800 System Sample Preparation Kit	240 Tests 960 Tests	P/N 05235782190 P/N 05235804190
cobas [®] 4800 System Lysis Kit 1	240 Tests 960 Tests	P/N 06768253190 P/N 06768270190
cobas [®] 4800 System Wash Buffer Kit	240 Tests 960 Tests	P/N 05235863190 P/N 05235871190
cobas [®] 4800 System Internal Control Kit 1	20 Runs	P/N 06768318190
cobas [®] 4800 Cdiff Amplification/Detection Kit	80 Tests	P/N 06768237190
cobas [®] 4800 Cdiff Controls and Cofactor Kit	10 Runs	P/N 06768300190

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Intended use

The **cobas**[®] Cdiff Test on the **cobas**[®] 4800 System is an automated, qualitative in vitro diagnostic test, that utilizes real-time polymerase chain reaction (PCR), for the direct detection of the toxin B (*tcd*B) gene of toxigenic *Clostridiodes difficile (C. difficile*) in unformed (liquid or soft) stool specimens obtained from patients suspected of having *C. difficile* infection (CDI). The **cobas**[®] Cdiff Test is intended for use as an aid in the diagnosis of CDI in humans in conjunction with clinical and epidemiological risk factors.

Summary and explanation of the test /principle of the procedure

Background: Screening of C. difficile

C. difficile is a Gram-positive, anaerobic, spore-forming bacillus that was identified as an etiologic agent of antibiotic-associated pseudomembranous colitis in the late 1970s.^{1,2} It is believed to be responsible for 15-20% of antibiotic-related cases of diarrhea and nearly all cases of antibiotic-associated pseudomembranous colitis.³ Over the last decade, the incidence of *C. difficile*-associated infection (CDI) has progressively increased and is now a significant clinical problem in developed countries. Whereas incidence rates ranged from 30-40 cases per 100,000 population in acute care hospitals in the United States, the incidence rose to more than 80 per 100,000 in 2005.⁴ Outbreaks of CDI have been previously described.⁵ The direct cost associated with CDI was \$6,326 per case in the US.⁶

Increases in incidence have in part been attributed to the emergence of a purportedly hypervirulent strain, classified as ribotype 027/North American pulsotype 1 (NAP1) and toxinotype III. Toxigenic strains of *C. difficile* typically produce two toxins: toxin A (an enterotoxin) and toxin B (a cytotoxin).⁷ A small percentage of strains only produce toxin B.⁸

Increased virulence has recently been described in strains that produce another toxin, termed binary toxin, and carry a deletion in the negative regulator gene tcdC.^{9,10} The latter strains were reported to be more virulent in vitro and appear to cause more morbidity and mortality in humans.^{11,12}

Following colonization with toxigenic *C. difficile*, individuals may become asymptomatic carriers or develop colonic disease. Clinical features of CDI may range from mild diarrhea to life-threatening pseudomembranous colitis characterized by abdominal pain, profuse diarrhea, and systemic symptoms such as fever, anorexia, nausea, and malaise.

Diagnosis of CDI is usually established by demonstration of the presence of toxins A and/or B in stool samples. Demonstration of the cytopathic effect on a monolayer of cells, by the action of toxin B, is considered by many to be the "gold standard."^{13, 14} Demonstration of the cytopathic effect can be achieved by direct incubation of stool supernatant on the monolayer of cells; alternatively, *C. difficile* isolates can be grown in selective broth and the supernatant obtained for subsequent incubation on the cell monolayer (toxigenic culture).¹⁵ Both techniques require a minimum of 48 to 72 hours for a final result. Immunoassays for the detection of toxins A and B are widely used because they provide positive results in less than 4 hours; however, sensitivities are significantly lower compared to tissue culture.¹⁶ Compared with clinical criteria supporting CDI, PCR was reported to have a sensitivity, specificity, and positive and negative predictive values of 93.3%, 97.4%, 75.5%, and 99.4%, respectively, with a turnaround time of <4 hours.¹⁵

PCR is considered the optimum rapid single test for detection of *C. difficile* toxin.¹⁷⁻²⁰ Despite the dramatic increase in incidence and severity of CDI, metronidazole or vancomycin remain the medical treatment of choice for acute episodes and recurrent infection.²¹

Infection control measures include the prudent use of antimicrobials, prevention of cross-infection, and active surveillance of cases.²² Repeat "test of cure" testing is not advised since toxins may be present for prolonged times without clinical symptoms.

Thus, there is a great need for highly sensitive and rapid automated detection of *C. difficile*. Molecular methods offer the potential to significantly reduce the detection time, thereby enabling the prompt initiation of antimicrobial treatment and the prompt implementation of infection control measures.¹⁷⁻²⁰

Explanation of the test

The **cobas**[®] Cdiff Test contains two major processes: (1) automated sample preparation to extract nucleic acids from the unformed stool specimens; (2) PCR amplification of target DNA sequences using *C. difficile* specific primers, and real-time detection of cleaved fluorescent-labeled *C. difficile* specific oligonucleotide detection probes. An Internal Control, containing unrelated randomized DNA sequence, is added to all samples prior to automated sample preparation and is amplified and detected simultaneously with each sample to monitor the entire process.

Principles of the procedure

Sample preparation

Sample preparation for the **cobas**[®] Cdiff Test is automated with the use of the **cobas**[®] x 480 instrument. Organisms are lysed with chaotropic agent, proteinase K, and SDS reagents. Released nucleic acids, along with added Internal Control DNA, are bound by magnetic glass particles. They are washed and then eluted into a small volume of buffer. The instrument then takes an aliquot of the eluted material and sets up the PCR reaction with an activated Master Mix.

PCR amplification and TaqMan[®] detection

The PCR cycling steps and detection of target signal occurs in the **cobas**[®] z 480 analyzer. The Master Mix reagent contains primer pairs and probes for two targets: toxin B and Internal Control. If the target nucleic acid sequences are present, amplification with the corresponding primers will occur by a thermostable DNA polymerase, generating PCR products (amplicon). These products are detected by specific TaqMan probes containing a fluorescent dye and a quencher. Normally, the quencher suppresses the fluorescence of the dye. However, if the PCR product is present, the probe hybridizes to the product and gets cleaved by the 5' to 3' nuclease activity of the polymerase. This reaction allows the fluorescence to be emitted from the dye, and the signal is recorded in real time during each PCR cycle by the **cobas**[®] z 480 analyzer. The signal is interpreted by the **cobas**[®] 4800 System Software and reported as final results.

Selective amplification

Selective amplification of target nucleic acid from the specimen is achieved in the **cobas**[®] Cdiff Test by the use of AmpErase (uracil-N-glycosylase) enzyme and deoxyuridine triphosphate (dUTP). The AmpErase enzyme recognizes and catalyzes the destruction of DNA strands containing deoxyuridine²³, but not DNA containing deoxythymidine. Deoxyuridine is not present in naturally occurring DNA, but is always present in amplicon due to the use of deoxyuridine triphosphate in place of thymidine triphosphate as one of the dNTPs in the Master Mix reagent; therefore, only amplicon contain deoxyuridine. Deoxyuridine renders contaminating amplicon susceptible to destruction by AmpErase enzyme prior to amplification of the target DNA. AmpErase enzyme, which is included in the Master Mix reagent, catalyzes the cleavage of deoxyuridine-containing DNA at the deoxyuridine residues by opening the deoxyribose chain at the C1-position. When heated in the first thermal cycling step at the alkaline pH of Master Mix, the amplicon DNA chain breaks at the position of the deoxyuridine, thereby rendering the DNA non-amplifiable. AmpErase enzyme is inactive at temperatures above 55°C, i.e., throughout the thermal cycling steps, and therefore does not destroy target amplicon. The **cobas**[®] Cdiff Test has been demonstrated to inactivate at least 1000 copies of deoxyuridine-containing *C. difficile* amplicon per PCR.

Materials, reagents, and specimen

Kit/Cassettes	Components and Reagent Ingredients	Quantity per Test	Safety Symbol and Warning
cobas [®] 4800 System Sample Preparation Kit 240 Tests (P/N: 05235782190)	MGP (cobas [®] 4800 System Magnetic Glass Particles) Magnetic Glass Particles 93% Isopropanol	10 x 4.5 mL	 DANGER H225 Highly flammable liquid and vapour. H319 Causes serious eye irritation. H336 May cause drowsiness or dizziness. P210 Keep away from heat, hot surfaces, sparks, open flames and other ignition sources. No smoking. P233 Keep container tightly closed. P261 Avoid breathing dust/ fume/ gas/ mist/ vapours/ spray. P280 Wear protective gloves/ eye protection/ face protection. P303 + P361 + P353 IF ON SKIN (or hair): Take off immediately all contaminated clothing. Rinse skin with water/shower. P370 + P378 In case of fire: Use dry sand, dry chemical or alcohol-resistant foam to extinguish.
	EB (cobas [®] 4800 System Elution Buffer)	10 x 18 mL	N/A
	Tris buffer 0.09% Sodium azide		

Materials and reagents provided

Kit/Cassettes	Components and Reagent Ingredients	Quantity per Test	Safety Symbol and Warning
cobas [®] 4800 System Sample Preparation Kit 960 Tests (P/N: 05235804190)	MGP (cobas [®] 4800 System Magnetic Glass Particles) Magnetic Glass Particles 93% Isopropanol	10 x 13.5 mL	 DANGER H225 Highly flammable liquid and vapour. H319 Causes serious eye irritation. H336 May cause drowsiness or dizziness. P210 Keep away from heat, hot surfaces, sparks, open flames and other ignition sources. No smoking. P233 Keep container tightly closed. P261 Avoid breathing dust/ fume/ gas/ mist/ vapours/ spray. P280 Wear protective gloves/ eye protection/ face protection. P303 + P361 + P353 IF ON SKIN (or hair): Take off immediately all contaminated clothing. Rinse skin with water/shower. P370 + P378 In case of fire: Use dry sand, dry chemical or alcohol-resistant foam to extinguish.
	EB		N/A
	(cobas [®] 4800 System Elution Buffer) Tris buffer 0.09% Sodium azide	10 x 18 mL	

Kit/Cassettes	Components and Reagent Ingredients	Quantity per Test	Safety Symbol and Warning
cobas [®] 4800 System Lysis Kit 1 240 Tests (P/N: 06768253190)	LYS-1 (cobas [®] 4800 System Lysis Buffer-1) Sodium citrate 5% Polydocanol 42.6% Guanidinium thiocyanate Dithiothreitol	10 x 10 mL	 DANGER H302: Harmful if swallowed. H314: Causes severe skin burns and eye damage. H411: Toxic to aquatic life with long lasting effects. EUH032: Contact with acids liberates very toxic gas. EUH071: Corrosive to the respiratory tract. P273: Avoid release to the environment. P280 Wear protective gloves/protective clothing/ eye protection/face protection/hearing protection. P303 + P361 + P353 IF ON SKIN (or hair): Take off immediately all contaminated clothing. Rinse skin with water. P304 + P340 + P310 IF INHALED: Remove person to fresh air and keep comfortable for breathing. Immediately call a POISON CENTER/doctor. P305 + P351 + P338 + P310 IF IN EYES: Rinse cautiously with water for several minutes. Remove contact lenses, if present and easy to do. Continue rinsing. Immediately call a POISON CENTER/ doctor. P391 Collect spillage. S93-84-0 Guanidinium thiocyanate 9002-92-0 Polidocanol 3483-12-3 (R*,R*)-1,4-dimercaptobutane-2,3-diol
	PK (cobas [®] 4800 System Proteinase K) Tris buffer EDTA Calcium chloride Calcium acetate < 2.0% Proteinase K Glycerine	10 x 0.9 mL	 DANGER H317: May cause an allergic skin reaction. H334: May cause allergy or asthma symptoms or breathing difficulties if inhaled. P261: Avoid breathing mist or vapours. P280: Wear protective gloves. P284: Wear respiratory protection. P304 + P340 IF INHALED: Remove person to fresh air and keep comfortable for breathing. P333 + P313 If skin irritation or rash occurs: Get medical advice/ attention. P342 + P311 If experiencing respiratory symptoms: Call a POISON CENTER/ doctor. 39450-01-6 Proteinase, Tritirachium album serine

Kit/Cassettes	Components and Reagent Ingredients	Quantity per Test	Safety Symbol and Warning
	SDS (cobas [®] 4800 System SDS Reagent) Tris buffer Sodium dodecyl sulfate 0.09% Sodium azide	10 x 3 mL	N/A
cobas [®] 4800 System Lysis Kit 1 960 Tests (P/N: 06768270190)	LYS-1 (cobas [®] 4800 System Lysis Buffer-1) Sodium citrate 5% Polydocanol 42.6% Guanidinium thiocyanate Dithiothreitol	10 x 36 mL	DANGERH302: Harmful if swallowed.H314: Causes severe skin burns and eye damage.H411: Toxic to aquatic life with long lasting effects.EUH032: Contact with acids liberates very toxic gas.EUH071: Corrosive to the respiratory tract.P273: Avoid release to the environment.P280: Wear protective gloves/ protective clothing/ eyeprotection/face protection/ hearing protection.P303 + P361 + P353 IF ON SKIN (or hair): Take offimmediately all contaminated clothing. Rinse skin with water.P304 + P340 + P310 IF INHALED: Remove person to fresh airand keep comfortable for breathing. Immediately call aPOISON CENTER/ doctor.P305 + P351 + P338 + P310 IF IN EYES: Rinse cautiouslywith water for several minutes. Remove contact lenses, ifpresent and easy to do. Continue rinsing. Immediately call aPOISON CENTER/ doctor.P391: Collect spillage.593-84-0 Guanidinium thiocyanate9002-92-0 Polidocanol3483-12-3 (R*,R*)-1,4-dimercaptobutane-2,3-diol

Kit/Cassettes	Components and Reagent Ingredients	Quantity per Test	Safety Symbol and Warning
	PK (cobas [®] 4800 System Proteinase K) Tris buffer EDTA Calcium chloride Calcium acetate < 2.0% Proteinase K Glycerine	20 x 1.2 mL	 DANGER H317: May cause an allergic skin reaction. H334: May cause allergy or asthma symptoms or breathing difficulties if inhaled. P261: Avoid breathing mist or vapours. P280: Wear protective gloves. P284: Wear respiratory protection. P304 + P340 IF INHALED: Remove person to fresh air and keep comfortable for breathing. P333 + P313 If skin irritation or rash occurs: Get medical advice/ attention. P342 + P311 If experiencing respiratory symptoms: Call a POISON CENTER/ doctor. 39450-01 - 6 Proteinase, Tritirachium album serine
	SDS (cobas [®] 4800 System SDS Reagent) Tris buffer Sodium dodecyl sulfate 0.09% Sodium azide	10 x 9 mL	N/A

Kit/Cassettes	Components and Reagent Ingredients	Quantity per Test	Safety Symbol and Warning
cobas [®] 4800 System Wash Buffer Kit 240 Tests (P/N: 05235863190)	WB (cobas [®] 4800 System Wash Buffer) Sodium citrate dihydrate 0.05% N-Methylisothiazolone HCl	10 x 55 mL	N/A
cobas [®] 4800 System Wash Buffer Kit 960 Tests (P/N: 05235871190)	WB (cobas [®] 4800 System Wash Buffer) Sodium citrate dihydrate 0.05% N-Methylisothiazolone HCl	10 x 200 mL	N/A
cobas [®] 4800 System Internal Control Kit 1 20 Runs (P/N: 06768318190)	IC-1 (cobas [®] 4800 IC-1) Tris buffer EDTA < 0.01% Poly rA RNA (synthetic) 0.05% Sodium azide < 0.01% Non-infectious, synthetic internal control DNA encapsulated in Lambda bacteriophage coat protein	20 x 0.5 mL	N/A
cobas [®] 4800 Cdiff Amplification/Detection Kit 80 Tests (P/N: 06768237190)	Cdiff MMX (cobas [®] Cdiff Master Mix) Tricine buffer EDTA DMSO Potassium acetate Potassium hydroxide Tween 20 < 0.19% dATP, dCTP, dGTP, dUTP < 0.01% Upstream and downstream <i>C. difficile</i> and Internal Control primers < 0.01% Fluorescent-labeled <i>C. difficile</i> and Internal Control probes < 0.01% Oligonucleotide aptamer < 0.01% Z05 DNA polymerase (microbial) < 0.02% AmpErase (uracil-N- glycosylase) enzyme (microbial) 0.09% Sodium azide	10 x 0.3 mL	N/A

Kit/Cassettes	Components and Reagent Ingredients	Quantity per Test	Safety Symbol and Warning
	Cdiff (+) C (cobas [®] Cdiff Positive Control) Tris buffer EDTA < 0.01% Poly rA RNA (synthetic) 0.05% Sodium azide < 0.01% Non-infectious plasmid DNA (microbial) containing <i>C.</i> <i>difficile</i> sequence	10 x 0.5 mL	N/A
cobas [®] 4800 Cdiff Controls and Cofactor Kit 10 Runs (P/N: 06768300190)	(-) C (cobas [®] 4800 System Negative Control) Tris buffer EDTA 0.05% Sodium azide < 0.01% Poly rA RNA (synthetic)	10 x 0.5 mL	N/A
	Cofactor-3 (cobas [®] 4800 Cofactor-3) Manganese acetate Magnesium acetate Bovine serum albumin from bovine plasma sourced in the United States 0.09% Sodium azide	10 x 1.7 mL	N/A

Reagent storage and handling

Reagent	Storage Temperature	Storage Time
cobas [®] 4800 System Sample Preparation Kit	2-8°C	Stable until the expiration date indicated
cobas [®] 4800 System Lysis Kit 1	2-8°C	Stable until the expiration date indicated
cobas [®] 4800 System Internal Control Kit 1	2-8°C	Stable until the expiration date indicated
cobas [®] 4800 Cdiff Amplification/Detection Kit	2-8°C	Stable until the expiration date indicated
cobas [®] 4800 Cdiff Controls and Cofactor Kit	2-8°C	Stable until the expiration date indicated
cobas® 4800 System Wash Buffer Kit	15-25°C	Stable until the expiration date indicated

Note: Do not freeze reagents.

Reagent expiry date is based on the Coordinated Universal Time (UTC). Local time for reagent expiry could be offset by plus or minus 12 hours, depending on the local time zone relative to UTC.

Additional materials required

Materials	P/N
CORE Tips, 1000 µL, rack of 96	04639642001
50 mL Reagent Reservoir	05232732001
200 mL Reagent Reservoir	05232759001
cobas [®] 4800 System Extraction (deep well) Plate	05232716001
cobas [®] 4800 System AD (microwell) Plate 0.3 mL and Sealing Film	05232724001
Sealing foil applicator	04900383001
24-position carrier	04639502001
Solid waste bag	05530873001 (small) or 04691989001 (large)
Hamilton STAR Plastic Chute	04639669001
cobas® PCR Media and Swab Sample Kit	07051891190
cobas [®] PCR Media Uni Swab Sample Kit	07958030190
Disposable gloves, powderless	Any powderless disposable gloves are acceptable.
Vortex Mixer (single tube)	Any vortex mixer is acceptable.
Centrifuge equipped with a swinging bucket rotor with minimum RCF of 1500	Any appropriate centrifuge is acceptable.

For more information regarding the materials sold separately, contact your local Roche representative.

Optional materials

Materials	P/N
Sealing mat or deep well plate cover	Roche 04789288001 or Hamilton 6474-01
Caps, neutral color (for recapping post-run specimens)	Roche P/N 07958056190; for recapping post-run specimens in 13 mL Round Base tubes.

For more information regarding the optional materials, contact your local Roche representative.

Instrumentation and software required but not provided

Required Instrumentation and Software, Not Provided
cobas [®] 4800 System
cobas® x 480 instrument
cobas® z 480 analyzer
Control Unit
cobas [®] 4800 System cobas [®] Cdiff AP Software Version 1.0.1 or higher

cobas® 4800 System Application Software (Core) Version 2.2.0 or higher

For more information regarding the materials sold separately, contact your local Roche representative.

Precautions and handling requirements

Warnings and precautions

As with any test procedure, good laboratory practice is essential to the proper performance of this assay. Due to the high analytical sensitivity of this test, care should be taken to keep reagents, specimens and amplification mixtures free of contamination.

- For in vitro diagnostic use only.
- Avoid microbial and DNA contamination of reagents and specimens.
- Safety Data Sheets (SDS) are available upon request from your local Roche office.
- LYS-1 reagent contains guanidine thiocyanate. Do not allow direct contact between guanidine thiocyanate and sodium hypochlorite (bleach) or other highly reactive reagents such as acids or bases. These mixtures can release a noxious gas.
- MGP contains isopropanol and is highly flammable. Keep away from open flames and potential spark producing environments.
- Prevent exposure of MGP to sources of magnetic field.
- EB, Cdiff MMX, SDS, Cofactor-3, (-)C, Cdiff (+)C and IC-1 contain sodium azide.
- For additional warnings, precautions and procedures to reduce the risk of contamination for the cobas[®] x 480 instrument or cobas[®] z 480 analyzer, consult the cobas[®] 4800 System User Assistance. If contamination is suspected, perform cleaning and weekly maintenance as described in the cobas[®] 4800 System User Assistance.
- Inform your local competent authority and manufacturer about any serious incidents which may occur when using this assay

Note: For specific instructions, see "Specimen collection, transport, and storage".

Good laboratory practice

- Do not pipette by mouth.
- Do not eat, drink or smoke in laboratory work areas.
- Wash hands thoroughly after handling specimens and kit reagents.
- Wear eye protection, laboratory coats and disposable gloves when handling any reagents. Avoid contact of these materials with the skin, eyes or mucous membranes. If contact does occur, immediately wash with large amounts of water. Burns can occur if left untreated. If spills occur, dilute with water before wiping dry.
- Thoroughly clean and disinfect all laboratory work surfaces with a freshly prepared solution of 0.5% sodium hypochlorite in distilled or deionized water (dilute household bleach 1:10). Follow by wiping the surface with 70% ethanol.

Contamination

- Gloves must be worn and must be changed between handling specimens and cobas[®] Cdiff reagents to prevent contamination. Avoid contaminating gloves when handling specimens and controls. Wear lab gloves, laboratory coats, and eye protection when handling specimens and kit reagents.
- Avoid microbial and ribonuclease contamination of reagents.
- False positive results may occur if carryover of specimens is not prevented during specimen handling.
- Specimens should be handled as infectious using safe laboratory procedures such as those outlined in *Biosafety in Microbiological and Biomedical Laboratories*²⁴ and in the CLSI Document M29-A4.²⁵

Integrity

- Do not use kits after their expiration dates.
- Do not pool reagents.
- Do not use disposable items beyond their expiration date.
- Do not use reagents or containers that are visibly damaged or show signs of leakage.
- All disposable items are for one time use. Do not reuse.
- All equipment should be properly maintained according to the manufacturer's instructions.

Disposal

cobas[®] 4800 system reagents and cobas[®] Cdiff Test specific reagents contain sodium azide (see "Warnings and precautions"). Sodium azide may react with lead and copper plumbing to form highly explosive metal azides. While disposing of solutions containing sodium azide down laboratory sinks, flush the drains with a large volume of cold water to prevent azide buildup.

• Dispose of unused reagents and waste in accordance with country, federal, state and local regulations.

Note: For disposal of liquid waste, refer to the cobas[®] 4800 System - User Assistance.

Note: Spillage and cleaning

- LYS-1 reagent contains guanidine thiocyanate. If liquid containing guanidine thiocyanate is spilled, clean with suitable laboratory detergent and water. If the spilled liquid contains potentially infectious agents, FIRST clean the affected area with laboratory detergent and water, and then with 0.5% sodium hypochlorite.
- If spills occur on the appropriate cobas[®] 4800 instrument, follow the instructions in the cobas[®] 4800 System User Assistance to clean.
- Do not use sodium hypochlorite solution (bleach) for cleaning the cobas[®] x 480 instrument or cobas[®] z 480 analyzer. Clean the cobas[®] x 480 instrument or cobas[®] z 480 analyzer according to procedures described in the appropriate cobas[®] 4800 System User Assistance.

Specimen collection, transport, and storage

Note: Handle all specimens as if they are capable of transmitting infectious agents.

Specimen collection

Collect unformed stool specimen in a sterile container. Specimens should be collected following the procedure documented in your institution's standard operating procedures.

Specimen transport storage and stability

Unformed stool specimens are stable at 2-30°C for 2 days, or 2-8°C for 7 days, and at -20°C for 60 days before testing on the **cobas**[®] 4800 System (this was demonstrated by testing specimens after consecutive storage at 30°C \pm 1°C for 2 days, followed by 2-8°C for 5 days, followed by -20°C for 60 days).

Stool specimen mixed with the **cobas**[®] PCR Media is stable at 2-8°C for 60 days or at 30°C for 7 days before testing on the **cobas**[®] 4800 System.

Transportation of *C. difficile* specimens must comply with country, federal, state and local regulations for the transport of etiologic agents.

Test procedure

Running the test

Workflow

Figure 1: cobas[®] Cdiff workflow

1	Start up the system.
2	Perform instrument maintenance.
3	Remove samples and reagents from storage.
4	Start run: • Load carriers with samples.
5	With LIS: confirm work order Without LIS: create work order
6	Load consumables (deepwell plate, microwell plate, tip racks) and reagents
7	Start sample preparation run
8	Unload and seal microwell plate
9	Remove samples, used reagents, and deepwell plate.
10	Load microwell plate into analyzer
11	Review results
12	With LIS: send results to LIS
13	Unload analyzer

Instructions for use

All reagents except Cdiff MMX and Cofactor-3 must be at ambient temperature prior to loading on the **cobas**[®] x 480 instrument. The Cdiff MMX and Cofactor-3 reagents may be taken directly from 2-8°C storage as they will equilibrate to ambient temperature on board the **cobas**[®] x 480 instrument by the time they are used in the process.

Note: Refer to the cobas[®] 4800 System - User Assistance for detailed operating instructions. Run size

The **cobas**[®] 4800 System is designed to support mixed-batch runs between the **cobas**[®] MRSA/SA, **cobas**[®] Cdiff and **cobas**[®] HSV 1 and 2 tests. The generic **cobas**[®] 4800 System Sample Preparation Kit, generic **cobas**[®] 4800 System Lysis Kit 1 and generic **cobas**[®] 4800 System Wash Buffer Kit are available in two kit sizes, each sufficient for 10 runs of up to either 24 or 96 samples, which include the controls and specimens for all assays to be run. The **cobas**[®] 4800 Cdiff Amplification/Detection Kit is sufficient to test up to 80 samples, which include Cdiff controls and specimens to be run. Multiple vials of the **cobas**[®] 4800 Cdiff Master Mix reagent can be used as appropriate in one run, as long as they are the same kit size. The generic **cobas**[®] 4800 System Internal Control Kit 1 and the **cobas**[®] 4800 Cdiff Controls and Cofactor Kit are available in a single kit size, and can support all run configurations. For each run containing *C. difficile* 06979394001-08EN specimens, one **cobas**[®] 4800 Cdiff Positive Control and one **cobas**[®] 4800 System Negative Control must be used (see "**Quality Control**"). For a single test run, the maximum number of samples allowed is 94 specimens and 2 controls.

Note: Although not an optimal use of reagents, a generic 96-Test reagent can be used for a run containing 1-22 total specimens. However, different sizes of the cobas[®] 4800 System Wash Buffer (WB) Kit, cobas[®] 4800 System Sample Preparation Kit and cobas[®] 4800 System Lysis Kit 1 cannot be mixed. For example, if a 96-Test WB reagent bottle is scanned at the start of the run, 96-Test size reagents from the other two kits must also be used.

Workflow

The **cobas**[®] Cdiff Test is performed using the full workflow within the **cobas**[®] 4800 Software. It consists of sample preparation on the **cobas**[®] x 480 instrument followed by amplification/detection on the **cobas**[®] z 480 analyzer. The run can be Cdiff only, or mixed-batched with the **cobas**[®] MRSA/SA and/or **cobas**[®] HSV 1 and 2 tests. Refer to the **cobas**[®] 4800 System - User Assistance for details.

Specimen transfer into cobas[®] PCR media tube

1. Use <u>one</u> polyester swab to transfer stool to the **cobas**[®] PCR Media tube – discard the second swab packaged in the packet (if applicable). Without touching the side of the stool container, immerse the tip of the swab fully into the stool specimen, up to the end of the tapered section, then promptly remove and place inoculated swab into the **cobas**[®] PCR Media tube. Do not test the sample if there is not enough stool to fully submerge the tip of the swab.

2. Break the swab shaft at the gray notch mark, by applying pressure against the side of the tube. Cap tube and vortex for a minimum of 5 seconds. Uncap and place tube(s) on 24-position sample carrier rack(s) for processing. Discard the caps.

- Note: The cobas[®] Cdiff Test has been validated for use with the cobas[®] PCR Media Kit, cobas[®] PCR Media Uni Swab Sample Kit, and cobas[®] PCR Media Dual Swab Sample Kit. Do not use other devices or media types.
- Note: Use only one polyester swab to transfer stool. Excess stool transferred to the cobas[®] PCR Media tube may cause clots and/or invalid results.
- Note: Stool specimens must be transferred into cobas[®] PCR Media tubes which are labeled with a proper barcode for processing on the cobas[®] x 480 instrument. Consult the cobas[®] 4800 System - User Assistance for proper barcoding procedures and the list of acceptable barcodes for the cobas[®] 4800 System.
- Note: To avoid cross-contamination of stool specimen suspensions in cobas[®] PCR Media, additional caps for cobas[®] PCR Media container in an alternate color (neutral; see "Optional materials") should be used to recap specimens after processing.

cobas[®] PCR Media contains sufficient volume for the stool suspension to be assayed multiple times on the cobas[®] 4800 System. Minimum stool suspension volume to conduct a cobas[®] Cdiff run is 3 mL in the cobas[®] PCR Media tube.

Performing the cobas[®] Cdiff Test

Note: Mixed batch runs between cobas[®] MRSA/SA and cobas[®] Cdiff and/or cobas[®] HSV 1 and 2 tests can be conducted. Refer to the cobas[®] 4800 System - User Assistance for more information.

- Perform the system startup and maintenance procedures by following the instructions in the **cobas**[®] 4800 System - User Assistance.
- 2. Collect all reagents and consumables needed. Reagents must be at room temperature by the time the run is started with the exception of **cobas**[®] Cdiff MMX and Cofactor-3 reagents.

Note: All reagents and reagent reservoirs are barcoded and designed for one time use. The cobas[®] 4800 Software tracks the use of the reagents and reagent reservoirs and rejects previously used reagents or reagent reservoirs.

- 3. Start a new run and define the work order for the run. There are three ways to create a work order:
 - By using the sample editor before sample rack is loaded into **cobas**[®] x 480 instrument ("Editor" button on the right of the main menu). Work orders can be saved, edited and reloaded if necessary.
 - By following the software wizard for the new run and loading specimens into **cobas**[®] x 480 instrument when prompted. The specimen barcodes will be automatically scanned, and the requested results for each specimen must be defined.
 - By using your institution's LIS system.

Refer to the **cobas**[®] 4800 System - User Assistance for more details. When selecting the requested results, check "Cdiff".

- 4. Load samples and define/select workorder or use LIS as appropriate. The "Unload sample carriers after transferring to deep well plate" option is selected by default. This allows the operator to retrieve the remaining stool suspension specimens as soon as possible after they are aliquoted for processing by cobas[®] x 480 instrument. Stool suspension containers should be re-capped with fresh closure (see "Optional materials") if storage is needed.
- 5. Follow the software wizard guide and load consumables. Do not load or remove individual tips into a partially used tip rack, as the software tracks the number of tips that are left. If there are not enough tips for the run to be conducted, the software will alert the user.
- 6. Load the sample preparation reagents into the barcoded reagent reservoirs. The reagent reservoirs are available in two sizes: 200 mL and 50 mL. Follow the software wizard guide to select the correct reagent reservoir size. The reagent reservoir barcodes must face to the right of the carrier. Use the "scan-scan-pour-place" method to load sample preparation reagents:
 - Scan the reagent bottle barcode
 - Scan the reagent reservoir barcode
 - Pour the reagent into the reservoir
 - Place the filled reagent reservoir into the designated position on the reagent carrier

- Note: The cobas[®] 4800 System has an internal clock to monitor the length of time the reagents are on-board. Once the WB is scanned, 1 hour is allowed to complete the loading process and click on the Start button. A countdown timer is displayed on the Workplace Tab. The system will not allow the run to start if the on-board timer has expired.
- Note: To assure the accurate transfer of MGP, vortex or vigorously shake the MGP vial <u>immediately prior</u> to dispensing into the reagent reservoir.
- Load amplification/detection reagents (Cdiff MMX and Cofactor-3), Proteinase K (PK) and controls [Cdiff (+) C, IC and (-) C] directly onto the reagent carrier. In order to prevent contamination, it is required to change gloves after handling positive controls.
- Note: The software wizard will calculate the optimal number and size of cobas[®] Cdiff MMX reagent to use. This will be reflected in the "Kit size" column on the MMX and Cofactor loading screen. To use a different size of cobas[®] Cdiff MMX reagent, click the "Change kit size" button.
- 8. Start sample preparation by clicking on "Start Run".
- 9. After a successful sample preparation run, the "Sample Preparation results" button and the Unload button become available. If desired, select "Sample Preparation results" button to review the results then select "Unload" to unload the plate carriers. Alternatively, select "Unload" to unload the plate carrier without reviewing the results. See the **cobas**[®] 4800 System User Assistance.
- 10. Follow the instructions in the **cobas**[®] 4800 System User Assistance to seal the microwell plate, transport the plate to the **cobas**[®] z 480 analyzer and start the amplification and detection run.
- Note: The cobas[®] 4800 System has an internal clock to monitor the length of time after addition of the prepared samples to activated master mix. Amplification and detection should be started as soon as possible but no later than 90 minutes after the end of the cobas[®] x 480 instrument run. A countdown timer is displayed on the Workplace Tab. The system will abort the run if the timer has expired.
- 11. When the amplification and detection run is completed, unload the microwell plate from the **cobas**[®] z 480 analyzer.
- 12. Follow the instructions in the **cobas**[®] 4800 System User Assistance to review and accept results.

Results

Quality control and validity of results

One set of **cobas**[®] Cdiff Test Positive and Negative Controls are included in each run. For any run, valid results must be obtained for both the Positive and Negative Control for the **cobas**[®] 4800 Software to display the reportable **cobas**[®] Cdiff Test results from that run.

Positive control

The Cdiff (+) Control contains non-infectious DNA plasmids of *C. difficile*. The Cdiff (+) Control monitors nucleic acid extraction, amplification, and detection steps in a given run of the test. The Cdiff (+) Control result must be 'Valid'. If the Cdiff (+) Control results are consistently invalid, contact your local Roche office for technical assistance.

Negative control

The (-) Control result must be 'Valid'. If the (-) Control results are consistently invalid, contact your local Roche office for technical assistance.

Internal control

The Internal Control is a recombinant bacteriophage lambda that contains randomized sequences and targets for internal control-specific primers and probe. The Internal Control is added to all specimens and the Positive and Negative Controls during sample preparation on the **cobas**[®] x 480 instrument. The Internal Control monitors nucleic acid extraction, amplification, and detection steps for a given specimen. The Internal Control is also required for validation of the run controls.

Interpretation of results

Note: All assay and run validation is determined by the cobas[®] 4800 Software.

Note: A valid run may include both valid and invalid specimen results.

For a valid run, specimen results are interpreted as shown in Table 1.

Table 1: Interpretation of results of the cobas[®] Cdiff Test

cobas [®] Cdiff Test	Result Report and Interpretation
POS Cdiff	Cdiff Positive Specimen is positive for the presence of <i>C. difficile</i> DNA.
NEG Cdiff	Cdiff Negative* C. difficile DNA, if present, could not be detected.
Invalid	Invalid Result is Invalid. The original specimen should be re-tested to obtain valid result. Place a new cap on the tube containing the stool suspension which had the invalid result and vortex for a minimum of 5 seconds. Add 0.5 mL of the vortexed stool suspension to a new cobas [®] PCR Media tube containing media. Cap the diluted tube and vortex for a minimum of 5 seconds. Uncap and place diluted tube on 24-position sample carrier for processing.
Failed	No Result for Specimen Consult the cobas [®] 4800 System - User Assistance for instructions to review run flags and recommended actions. In rare cases when pipetting error (e.g. clot or other obstruction) occurs, the original stool specimen suspension tube should be closed with a new cap and placed into a centrifuge. Accelerate to 1800 RCF (or 1800 x g) then stop. Ensure the vial is not shaken or mixed after centrifuge. Uncap and place tube on 24-position sample carrier for processing.

* A negative result does not preclude the presence of *C. difficile* DNA because results depend on adequate specimen collection, absence of inhibitors, and sufficient DNA to be detected.

Invalid results may be obtained if the specimen contains excess stool or inhibitory substances that prevent nucleic acid target extraction and/or amplification and detection. See "**Procedural limitations**" for known interference substances.

Note: The minimum volume of stool suspension necessary for the **cobas**® Cdiff Test is 3 mL.

List of result flags

The following table lists flags which are relevant for result interpretation.

cobas [®] Cdiff Test	cobas [®] Cdiff Test	Result Report and Interpretation
R20	The positive control is invalid.	An external control is invalid.
		1. Repeat entire run with fresh reagents.
		2. If the problem persists, contact Roche Service.
R21	The negative control is invalid.	An external control is invalid.
		1. Repeat entire run with fresh reagents.
		2. If the problem persists, contact Roche Service.
Х3	Error: Clot was detected Sample was not processed.	Make sure that the samples were handled according to the workflow description.
		1. Check the sample for clots.
		2. Rerun the sample.
X4	Error: Pipetting error occurred. Sample was not processed.	Insufficient sample volume or mechanical error during pipetting is the most likely reason.
		1. Make sure that there is enough sample volume.
		2. Check whether the tip eject plate is placed correctly.
		3. Rerun the sample.

 Table 2:
 List of flags for the cobas[®] Cdiff Test

Procedural limitations

- 1. The **cobas**[®] Cdiff Test has only been validated for use with unformed stool specimens that have been transferred into the **cobas**[®] PCR Media according to this Instructions-For-Use document.
- Reliable results are dependent on adequate specimen collection, transport, storage, and processing. Follow the procedures in this Instructions-For-Use document (also referred to as a Package Insert) and the cobas[®] 4800 System - User Assistance.
- 3. Detection of *C. difficile* DNA is dependent on the number of organisms present in the specimen and may be affected by specimen collection/processing methods, history of hospitalization, antibiotic treatment regime, and *C. difficile* strains.
- 4. False negative or invalid results may occur due to interference from various substances. The Internal Control is included in the **cobas**[®] Cdiff Test to help identify the specimens containing substances that may interfere with nucleic acid isolation and PCR amplification. Known interference includes, but may not be limited to the following:
 - Specimens containing greater than 25% (w/v) mucin may generate false negative results.
- 5. A positive result is indicative of the presence of *C. difficile* DNA and not necessarily viable organisms. Therefore, a positive result does not necessarily mean eradication treatment failure.
- 6. Mutations or polymorphisms in primer- or probe-binding regions may affect detection of new or unknown variants, resulting in a false negative result with the **cobas**[®] Cdiff Test.
- 7. The predictive value of an assay depends on the prevalence of the disease in any particular population.
- The addition of AmpErase enzyme into the **cobas**[®] 4800 Cdiff Master Mix enables selective amplification of target DNA; however, good laboratory practices and careful adherence to the procedures specified in this Instructions-For-Use document are necessary to avoid contamination of reagents and amplification mixtures.
- 9. Use of this product must be limited to personnel trained in the techniques of PCR and the use of the **cobas**[®] 4800 System.
- 10. Only the **cobas**[®] x 480 instrument and **cobas**[®] z 480 analyzer have been validated for use with this product. No other sample preparation instrument or PCR System can be used with this product.
- 11. Due to inherent differences between technologies, it is recommended that, prior to switching from one technology to the next, users perform method correlation studies in their laboratory to identify technology differences, and verify the new procedure. One hundred percent agreement between the results should not be expected due to aforementioned differences between technologies.
- 12. Cross-contamination can cause false positive results. The **cobas**[®] 4800 System is an automated realtime PCR instrument designed to minimize the risk of cross-contamination during specimen processing, nucleic acid extraction, amplification and detection. To challenge the robustness of the system, a simulated, non-clinical checkerboard study was performed on a panel of alternating high positive and negative contrived samples, to assess a theoretical cross-contamination rate of the system. The high

positive samples had Ct values earlier than would be observed in 95% of infected patients in the intended use population. The cross-contamination rate in this checkerboard study was determined to be 0.24% (1/423). Cross-contamination rates in clinical settings depend on the proportion of high positive samples and prevalence of disease. Routine clinical cross-contamination rates are expected to be dramatically lower than what was observed in this study and need to be assessed in user's settings.

Non-clinical performance evaluation

Analytical sensitivity

The analytical sensitivity (Limit of Detection or LOD) for the **cobas**[®] Cdiff Test was determined by analyzing quantified *C. difficile* cultures diluted to multiple concentration levels in negative stool background suspension in **cobas**[®] PCR Media. All levels were tested using the **cobas**[®] Cdiff Test across three unique lots of **cobas**[®] Cdiff Test reagents. At least 21 replicates per reagent lot were tested at each level. LOD for this test is defined as the target concentration which can be detected as positive in \geq 95% of the replicates tested, based on results generated by the worst performing reagent lot.

The seven C. difficile strains tested in the analytical sensitivity study are shown in Table 3.

	_	REA*	PFG [†]		Phenotype	LOD (CFU/swab)	
Strain ID	Toxinotype	Туре	Туре	Ribotype		By Positive Rate	By Probit Analysis
ATCC 43255 (VPI 10463)	0	N/A	N/A	087	A+B+CDT-	113	90
ATCC BAA-1382 (630)	0	R 23	N/A	012	A+B+CDT-	81	83
CDC 204118	III	BI 8	NAP1	027	A+B+CDT+	54	42
R12087 (CD196)	III	BI	NAP1	027	A+B+CDT+	54	54
2748-06	V	N/A	N/A	078	N/A	54	45
ATCC 43598 (1470)	VIII	N/A	N/A	017	A-B+	225	130
F15	XII	N/A	N/A	N/A	N/A	54	59

 Table 3:
 cobas[®] Cdiff Test LOD (Limit of Detection)

* Restriction endonuclease analysis; [†]Pulse Field Gel

Detection of C. difficile genotypes

The limit of detection (LOD) of the **cobas**[®] Cdiff Test on 28 toxigenic strains representing additional toxinotypes were verified by testing 40 replicates per level at multiple levels. Dilutions and testing samples were prepared in a similar fashion as in the Limit of Detection study described above. The lowest level that had at least 95% observed hit rate are summarized in Table 4.

All 28 toxigenic strains (Table 4) were detected as positive in \geq 95% of the replicates tested at concentrations ranging from 77.9 CFU/swab to 460 CFU/swab.

Strain	Toxinotype	Ribotype	Conc. (CFU/swab)	Positive Rate
EX 623	I	102	77.9	95.0%
AC 008	II	103	77.9	95.0%
SE 844	Illa	080	234	100.0%
55767	IV	023	77.9	100.0%
SE 881	V	045	234	100.0%
51377	VI	N/A	234	100.0%
57267	VII	063	77.9	97.5%
51680	IX	019	77.9	100.0%
8864	Х	036	77.9	97.5%
R 9367	XIII	070	77.9	97.5%
R 10870	XIV	111	234	100.0%
R 9385	XV	122	234	100.0%
SUC36	XVI	078	234	100.0%
J9965	XVII	N/A	460	97.5%

 Table 4:
 Summary of toxigenic C. difficile verification results

 Table 4:
 Summary of Toxigenic C. difficile Verification Results (Continued)

Strain	Toxinotype	Ribotype	Conc. (CFU/swab)	Positive Rate
K095	XVIII	014	234	95.0%
TR13	XIX	N/A	234	97.5%
TR14	XX	N/A	77.9	100.0%
CH6223	XXI	N/A	234	100.0%
CD07-468	XXII	N/A	234	100.0%
8785	XXIII	N/A	234	95.0%
597B	XXIV	131	234	97.5%
7325	XXV	027	234	100.0%
7459	XXVI	N/A	234	95.0%
KK2443-2006	XXVII	N/A	234	100.0%
CD08-070	XXVIII	126	234	97.5%
CD07-140	XXIX	056	234	97.5%
ES 130	XXX	N/A	234	100.0%
WA 151	XXXI	N/A	460	100.0%

Precision

In-house precision study was conducted using a panel composed of *C. difficile* cultures diluted into negative stool suspension in **cobas**[®] PCR Media to concentration levels below Limit of Detection (LOD), near LOD and above LOD of the **cobas**[®] Cdiff Test. A negative level composed of only the stool suspension in **cobas**[®] PCR Media was also tested. The study used three unique lots of **cobas**[®] Cdiff Test reagents and three instruments for a total of 36 runs over 12 days. A description of the precision panels and the study summary is shown in Table 5. Analysis of the variance components (Table 6) suggested that most variability of target Ct values is attributed to within run (random) and lot to lot factors (60.0% and 25.3%, respectively) for concentration level at or around LOD. For concentration level above LOD, most of the Ct value variability is attributed to within run (random) and instrument to instrument factors (72.5% and 24.7%, respectively). Results (Table 7) show that the target Ct values had overall CV (%) of 1.5% for concentration level at LOD and 1.1% for concentration level above LOD.

Panel Member	N	N	Desitive Data	95% CL	
Panel Weinber	Tested	Positive	Positive Rate	Lower	Upper
Negative	72	0	0.0%	0.0%	5.0%
< 1 x LOD	72	21	29.2%	19.0%	41.1%
~ 1 x LOD	72	72	100.0%	95.0%	100.0%
~ 3 x LOD	72	72	100.0%	95.0%	100.0%

Table 5:In-house precision study positive rate analysis

LOD = Limit of Detection

Table 6: Variance components analysis for precision panel members

Laval	Mean	Va	Total				
Level	wean	Lot	Instrument	Kit Size	Day	Random	Total
~ 1 x LOD	38.5	0.0789	0.0189	0.0001	0.0270	0.1875	0.3123
~ 1 X LOD	30.0	25.25%	6.04%	0.03%	8.65%	60.03%	100.00%
	07 F	0.0047	0.0404	0.0000	0.0000	0.1188	0.1638
~ 3 x LOD	37.5	2.84%	24.65%	0.00%	0.00%	72.51%	100.00%

LOD = Limit of Detection

 Table 7:
 Standard deviations and coefficients of variation (%) analysis for precision panel members

			SD Components/ CV (%)					
Level	Mean	Lot	Instrumen t	Kit Size	Day	Random	Total	
	00 F	0.28	0.14	0.01	0.16	0.43	0.56	
~ 1 x LOD	38.5	0.73%	0.36%	0.03%	0.43%	1.12%	1.45%	
	07.5	0.07	0.20	0.00	0.00	0.34	0.40	
~ 3 x LOD	37.5	0.18%	0.54%	0.00%	0.00%	0.92%	1.08%	

LOD = Limit of Detection

Analytical specificity

To assess the analytical specificity of the **cobas**[®] Cdiff Test, the following organism panels were tested: 1) 103 bacteria, fungi and viruses that may be found in stool specimens and one human cell (Table 8); 2) 28 *Clostridiaceae* genus organisms, including non-toxigenic *C. difficile* (* Cytomegalovirus (HHV5) was spiked at 2.0 x 10³ PFU/mL, Human Adenovirus Type 40 at 2.2 x 10³ PFU/mL, and Human Rotavirus at 9.8 x 10³ PFU/mL for testing- Table 9).

All bacteria and human cells were spiked to 1×10^6 Units*/mL, and all viruses were spiked to 1×10^5 Units*/mL, except for Cytomegalovirus (HHV5), Human Adenovirus Type 40, and Human Rotavirus, which were spiked to lower concentrations due to stock concentration limitations. Testing was performed with the organisms alone or with two *C. difficile* isolates present individually at 3 x Limit of Detection (LOD) of the **cobas**[®] Cdiff Test. Results indicated that none of these organisms interfered with detection of intended Cdiff targets. None produced false positive results when there was no intended *C. difficile* target present.

Clostridium botulinum analytical specificity was confirmed using BLAST program against GenBank nucleotide sequence database to mimic PCR amplicon generation step.

*Bacteria were quantified as colony forming units (CFU)/mL, human cell were quantified as cells/mL, and viruses were quantified as plaque forming units (PFU)/mL, except following microorganisms. *Chlamydia trachomatis* was quantified as elementary body (EB)/mL. Cytomegalovirus, Human Echovirus, and Human Enterovirus were quantified as IU/mL.

Abiotrophia defectiva	Acinetobacter baumannii	Acinetobacter lwoffii	
Aeromonas hydrophila	Alcaligenes faecalis subsp. Faecalis	Anaerococcus tetradius	
Bacillus cereus ATCC 11778	Bacillus cereus ATCC 13472	Bacteroides caccae	
Bacteroides merdae	Bacteroides stercoris	Bifidobacterium adolescentis	
Bifidobacterium longum	Campylobacter coli	Campylobacter jejuni	
Candida albicans	Candida catenulata	Cedecea davisae	
Chlamydia Trachomatis Serovar L2	Citrobacter amalonaticus	Citrobacter freundii	
Citrobacter koseri	Citrobacter sedlakii	Collinsella aerofaciens	
Corynebacterium genitalium	Desulfovibrio piger	Edwardsiella tarda	
Eggerthella lenta	Enterobacter aerogenes	Enterobacter cloacae	
Enterococcus casseliflavus	Enterococcus cecorum	Enterococcus dispar	
Enterococcus faecalis	Enterococcus faecium vanA	Enterococcus gallinarum vanC	
Enterococcus hirae	Enterococcus raffinosus	Escherichia coli ATCC 11775	
Escherichia coli ATCC 25922	Escherichia fergusonii	Escherichia hermannii	
Fusobacterium varium	Gardnerella vaginalis	Gemella morbillorum	
Hafnia alvei	HCT-15 Human Cells	Helicobacter fennelliae	
Helicobacter pylori	Klebsiella oxytoca	Klebsiella pneumoniae subsp. pneumoniae	
Lactobacillus acidophilus	Lactobacillus reuteri	Lactococcus lactis	
Leminorella grimontii	Listeria grayi	Listeria innocua	
Listeria monocytogenes	Mitsuokella multacida	Mobilincus curtisii	
Moellerella wisconsensis	Morganella morganii	Neisseria gonorrhoeae	

 Table 8:
 Microorganisms and human cells

Peptoniphilus asaccharolyticus	Peptostreptococcus anaerobius	Plesiomonas shigelloides	
Porphyromonas asaccharolytica	Prevotella melaninogenica	Proteus mirabilis	
Proteus penneri	Providencia alcalifaciens	Providencia rettgeri	
Providencia stuartii	Pseudomonas aeruginosa	Pseudomonas putida	
Ruminococcus bromii	Salmonella choleraesuis subsp. choleraesuis	Salmonella enterica subsp. arizonae (f.k.a. Salmonella choleraesuis subsp. arizonae)	
Salmonella enterica subsp. enterica serovar Choleraesuis	Serratia liquefaciens	Serratia marcescens	
Shigella boydii	Shigella dysenteriae	Shigella sonnei	
Staphylococcus aureus	Staphylococcus epidermidis	Stenotrophomonas maltophilia	
Streptococcus agalactiae	Streptococcus dysgalactiae	Streptococcus intermedius	
Streptococcus uberis	Trabulsiella guamensis	Veillonella parvula	
Vibrio cholerae	Vibrio parahaemolyticus	Yersinia bercovieri	
Yersinia rohdei	Cytomegalovirus (HHV5)*	Human Adenovirus Type 40*	
Human Coxsackievirus A10	Human Enterovirus 11	Human Enterovirus 71	
Human Rotavirus*	Norovirus GII	-	

* Cytomegalovirus (HHV5) was spiked at 2.0 x 10³ PFU/mL, Human Adenovirus Type 40 at 2.2 x 10³ PFU/mL, and Human Rotavirus at 9.8 x 10³ PFU/mL for testing.

Table 9:	Clostridiaceae organisms,	including non-toxigenic C. difficile
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Clostridium beijerinckii	Clostridium bifermentans	Clostridium bolteae
Clostridium botulinum*	Clostridium butyricum	Clostridium chauvoei
Clostridioides difficile Serogroup B (non-toxigenic)	Clostridioides difficile Serogroup I (non-toxigenic)	Clostridium fallax
Clostridium haemolyticum	Clostridium histolyticum	Clostridium innocuum
Clostridium methylpentosum	Clostridium nexile	Clostridium novyi
Clostridium orbiscindens (re-named Flavonifractor plautii)	Clostridium paraputrificum	Clostridium perfringens
Clostridium ramosum	Clostridium scindens	Clostridium septicum
Clostridium sordellii	Clostridium sphenoides	Clostridium spiroforme
Clostridium sporogenes	Clostridium symbiosum	Clostridium tertium
Clostridium tetani	-	-

* Based on BLAST program.

Interference

Twenty-six commonly used medications, as well as fecal fat, whole blood, and mucin, were tested for potential interference effects with the **cobas**[®] Cdiff Test. All substances were tested at levels above what could be reasonably expected to be collected by a swab in a stool specimen. The amount of interference substance is expressed as concentration in primary stool specimen. Two *C. difficile* isolates were spiked to 3 x Limit of Detection (LOD) of the **cobas**[®] Cdiff Test and used as targets in the tests. No interference was observed for exogenous substances. For fecal fat, no interference was observed up to 28%, for whole blood, no interference was observed up to 50%, and for mucin, no interference was observed up to 25%. These results are summarized in Table 10.

Table 10:	Results from interference substances testing
Table IV.	Results nom interference substances testing

Substance	Concentration	Results
Fecal Fat	4 ~ 28 % (w/v)	No interference
Whole blood	25, 50 % (v/v)	No interference
Mucin	25, 50 % (w/v)	No interference up to 25% (w/v)
Tums	10% (w/v)	No interference
Vancomycin	1% (w/v)	No interference
Metronidazole	10% (w/v)	No interference
Imodium AD®	10% (w/v)	No interference
Stool Softener	10% (w/v)	No interference
Pepto-Bismol [®] (Procter & Gamble)	10% (v/v)	No interference
Nystatin Ointment USP	10% (w/v)	No interference
Preparation H [®] with Bio-Dyne [®] Cream (Wyeth)	10% (w/v)	No interference
GYNOL II	10% (w/v)	No interference
Vagisil [®] Anti-itch cream	10% (w/v)	No interference
Anusol [®] Plus	10% (w/v)	No interference
Sunscreen	1% (w/v)	No interference
Monistat [®] 7	10% (w/v)	No interference
Vaseline™	10% (w/v)	No interference
SAB-Dimenhydrinate [®] Suppositories (SABEX [®])	10% (w/v)	No interference
Mineral Oil	10% (v/v)	No interference
Equate Natural Vegetable Laxative	10% (w/v)	No interference
Dulcolax®	10% (w/v)	No interference
Fleet [®] (CB Fleet Company)	10% (w/v)	No interference
K-Y Jelly/Gelée [®] (McNeil-PPC)	1% (w/v)	No interference
Afrin Original Nasal Spray	10% (v/v)	No interference
Witch hazel	Liquid from 1 wipe/swab	No interference
E-Z-HD [™] High Density Barium Sulfate for suspension (E-Z-EM Canada)	20% (w/v)	No interference
Palmitic acid	10% (w/v)	No interference
Stearic acid	10% (w/v)	No interference
Aleve	10% (w/v)	No interference

Clinical performance using Clinical Specimens

The performance of the **cobas**[®] Cdiff Test was compared to a FDA-cleared and CE-marked State-of-the-Art comparator NAT, using tissue culture cytotoxicity testing on the *C. difficile* isolates from direct culture as reference method. Stool specimen collected at 3 hospitals/medical center were tested by **cobas**[®] Cdiff Test and comparator NAT and sent to a reference laboratory for tissue culture cytotoxicity testing.

The **cobas**[®] Cdiff Test and State-of-the-Art comparator NAT test were performed per the manufacturers' instructions. Tissue culture cytotoxicity test was performed using direct culture procedure. Each stool specimen was inoculated onto pre-reduced cycloserine-cefoxitin-fructose agar (CCFA-HT). Suspected

colonies were identified as *C. difficile* by Gram staining, aero-intolerance, and by the Pro-Disk Test and inoculated into anaerobic chopped meat broth. Supernatants obtained from anaerobic chopped meat broth will then be processed for the detection *C. difficile* toxin B using tissue culture cytotoxicity test (C. DIFFICILE TOX-B test, Techlab).

A total of 1,434 subjects were enrolled from five sites. One hundred fifty six subjects were excluded due to incomplete results. There were 152 *C. difficile* positive specimens by direct culture (prevalence: 11.9%). The performance of the **cobas**[®] Cdiff Test and the comparator NAT against direct culture is shown in Table 11 and Table 12.

Table 11:	cobas [®] Cdiff Test against direct culture
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		Direct Culture		
		Positive	Negative	Total
cobas [®] Cdiff Test	Positive	147	30	177
	Negative	5	1,096	1,101
	Total	152	1,126	1,278

Sensitivity and specificity

Sensitivity and Specificity of the **cobas**[®] Cdiff Test to direct culture are 96.7% (95% two-sided confidence interval: 92.5-98.6%) and 97.3% (95% two-sided confidence interval: 96.2-98.1%), respectively.

Positive and negative predictive value

Observed positive and negative predictive values of the **cobas**[®] Cdiff Test for study samples are 83.1% (95% two-sided confidence interval: 76.7-88.3%) and 99.5% (95% two-sided confidence interval: 98.9-99.9%), respectively.

Table 12:	Nucleic Acid Test (NAT) comparator against direct culture
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		Direct Culture		
		Positive	Negative	Total
NAT Comparator	Positive	147	29	176
	Negative	5	1,097	1,102
	Total	152	1,126	1,278

Sensitivity and specificity

Sensitivity and specificity of the NAT comparator to direct culture are 96.7% (95% two-sided confidence interval: 92.5-98.6%) and 97.4% (95% two-sided confidence interval: 96.3-98.2%), respectively.

Correlation to NAT comparator

The performance of the **cobas**[®] Cdiff Test in direct comparison to a FDA-cleared and CE-marked State-of-the-Art comparator NAT is shown in Table 13.

		Comparator NAT		
		Positive	Negative	Total
cobas [®] Cdiff Test	Positive	167	10	177
	Negative	9	1,092	1,101
	Total	176	1,102	1,278

Percent positive and negative agreements

Percent Positive and Negative Agreements of the **cobas**[®] Cdiff Test to NAT comparator are 94.9% (95% two-sided confidence interval: 90.6-97.3%) and 99.1% (95% two-sided confidence interval: 98.3-99.5%), respectively.

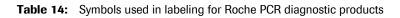
Additional information

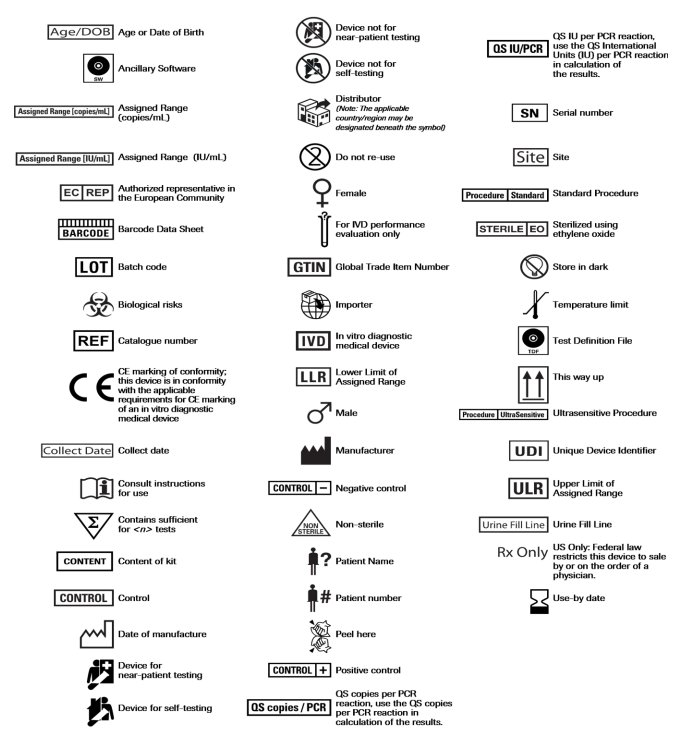
Key assay features

Sample type	Unformed stool specimens
Amount of sample required	4.3 mL of cobas[®] PCR media in the primary vial, a minimum of 3 mL is required for a cobas[®] Cdiff Test.
Test duration	Results are available within 2.5 hours after loading the specimen on the system. (1-22 specimens)
Analytical sensitivity	From 54 to 460 CFU/swab depending on isolate.
Specificity	No cross-reactivity with 125 closely related organisms or organisms typically found in stool specimens.
Inclusivity	All known <i>C. difficile</i> (Toxinotypes 0 ~ XXXI, except non-Toxigenic Toxinotypes XI) including the BI/ NAP1/027 hyper-virulent epidemic strain.

Symbols

The following symbols are used in labeling for Roche PCR diagnostic products.





Technical support

For technical support (assistance) please reach out to your local affiliate: https://www.roche.com/about/business/roche_worldwide.htm

Manufacturer and importer

Table 15: Manufacturer and importer



Roche Molecular Systems, Inc. 1080 US Highway 202 South Branchburg, NJ 08876 USA www.roche.com

Made in USA



Roche Diagnostics GmbH Sandhofer Strasse 116 68305 Mannheim Germany

Trademarks and patents

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Document revision

Document Revision Information	
Doc Rev 7.0 04/2023	Change Clostridium to Clostridiodes throughout the document Removed all references to Cdiff 240T test kit due to obsolescence Updated Precautions and handling requirements section to advise user to reach out to local competent authority. Added "damage/leakage" warning to Integrity section Renamed Correlation section to Clinical Performance using Clinical Specimens section. Added weblink to the summary of safety and performance report. Added CE mark symbol with notified body number. Updated the harmonized symbol page. Updated to current economic operators. Updated Trademarks and patents section, including the link. Inserted Rx Only symbol on first page. Added Made in statement. Added Technical support section. Updated cobas [®] branding. Please contact your local Roche Representative if you have any questions.
Doc Rev 8.0 02/2024	Updated Lysis Kits 1 hazard information. Updated cobas [®] branding. Please contact your local Roche Representative if you have any questions.

The summary of safety and performance report can be found using the following link: https://ec.europa.eu/tools/eudamed.