



Universiteit
Leiden
The Netherlands

Towards understanding *Clostridioides difficile* colonization

Crobach, M.J.T.

Citation

Crobach, M. J. T. (2024, February 14). *Towards understanding Clostridioides difficile* colonization. Retrieved from <https://hdl.handle.net/1887/3717585>

Version: Publisher's Version

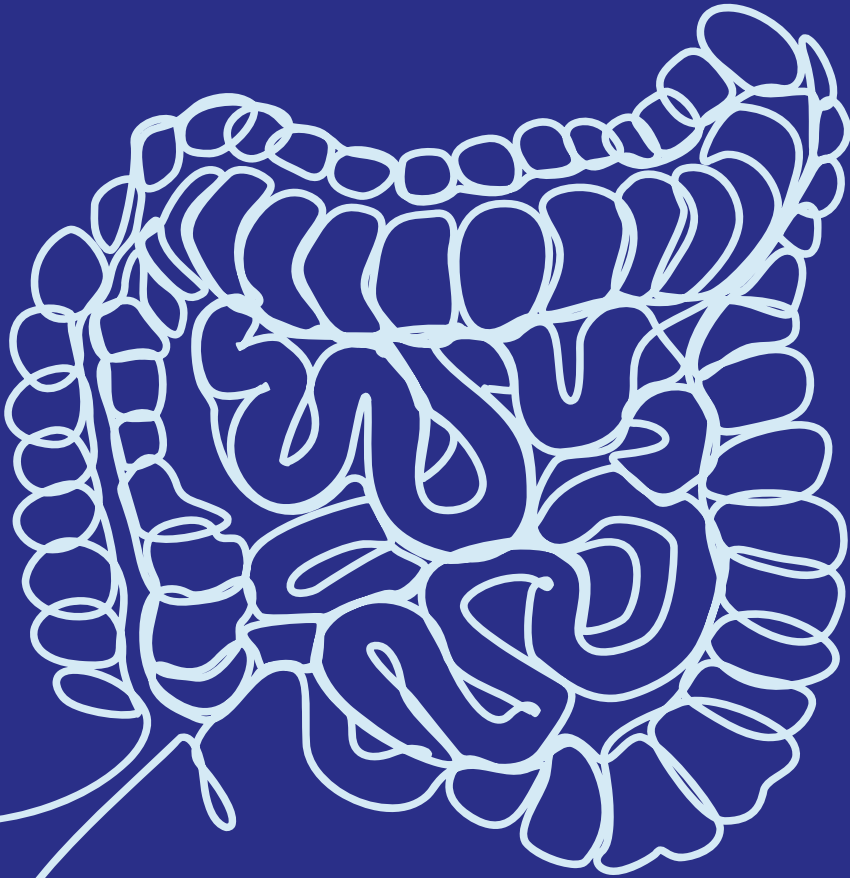
License: [Licence agreement concerning inclusion of doctoral thesis in the Institutional Repository of the University of Leiden](#)

Downloaded from: <https://hdl.handle.net/1887/3717585>

Note: To cite this publication please use the final published version (if applicable).

CHAPTER 3

European Society of Clinical Microbiology and Infectious Diseases: update of the diagnostic guidance document for *Clostridium difficile* infection



Clinical Microbiology and Infection, 2016

M.J.T. Crobach¹

T. Planche²

C. Eckert³

F. Barbut³

E.M. Terveer¹

O.M. Dekkers⁴

M.H. Wilcox⁵

E.J. Kuijper¹

¹ Department of Medical Microbiology, Centre for Infectious Diseases, Leiden University Medical Center, Leiden, The Netherlands

² Department of Medical Microbiology, St. George's Hospital, London, UK

³ National Reference Laboratory for *Clostridium difficile*, Paris, France

⁴ Departments of Clinical Epidemiology and Internal Medicine, Leiden University Medical Center, Leiden, The Netherlands and Department of Clinical Epidemiology, Aarhus University, Aarhus, Denmark

⁵ Department of Microbiology, Leeds Teaching Hospitals & University of Leeds, Leeds, UK

Clin Microbiol Infect 2016 Aug;22 Suppl 4:S63-81

Supplementary information available online



Abstract

In 2009 the first European Society of Clinical Microbiology and Infectious Diseases (ESCMID) guideline for diagnosing *Clostridium difficile* infection (CDI) was launched. Since then newer tests for diagnosing CDI have become available, especially nucleic acid amplification tests. The main objectives of this update of the guidance document are to summarize the currently available evidence concerning laboratory diagnosis of CDI and to formulate and revise recommendations to optimize CDI testing. This update is essential to improve the diagnosis of CDI and to improve uniformity in CDI diagnosis for surveillance purposes among Europe. An electronic search for literature concerning the laboratory diagnosis of CDI was performed. Studies evaluating a commercial laboratory test compared to a reference test were also included in a meta-analysis. The commercial tests that were evaluated included enzyme immunoassays (EIAs) detecting glutamate dehydrogenase, EIAs detecting toxins A and B and nucleic acid amplification tests. Recommendations were formulated by an executive committee, and the strength of recommendations and quality of evidence were graded using the Grades of Recommendation Assessment, Development and Evaluation (GRADE) system. No single commercial test can be used as a stand-alone test for diagnosing CDI as a result of inadequate positive predictive values at low CDI prevalence. Therefore, the use of a two-step algorithm is recommended. Samples without free toxin detected by toxins A and B EIA but with positive glutamate dehydrogenase EIA, nucleic acid amplification test or toxigenic culture results need clinical evaluation to discern CDI from asymptomatic carriage.

Introduction

The previous European Society of Clinical Microbiology and Infectious Diseases (ESCMID) guidance document for *Clostridium difficile* infection (CDI) was published in 2009. (1) Since then many laboratories in Europe have implemented a diagnostic algorithm for diagnosing CDI. However, many new diagnostic tests have become available in the meantime, especially nucleic acid amplification tests (NAATs). Although several of these tests have been marketed, their role in the diagnosis of CDI needs to be clarified. Also, the importance of free toxin detection in stool needs to be addressed. This update of the previous guidance document is essential to improve the diagnosis of CDI; to optimize its management, prevention and control; and to improve uniformity in CDI diagnosis for surveillance purposes across Europe.

The main objectives of this guidance document are to summarize the currently available evidence concerning laboratory diagnosis of CDI and to formulate recommendations to optimize CDI testing. This guideline is intended for use among medical microbiologists, gastroenterologists, infectious disease specialists and infection control practitioners. The target population is diarrhoeal patients suspected of having CDI.

Material and Methods

To be able to revise our previous recommendations, an update of the 2009 meta-analysis was performed. In addition, other guidelines and recent literature concerning the diagnosis of CDI were reviewed.

Update of meta-analysis

Search strategy

Studies evaluating laboratory assays for diagnosing CDI were searched in PubMed, Embase, Web of Science, Central and the Cochrane Library. Searches were performed in June 2014 with the support of a trained librarian. The search was restricted to articles published since 2009 in the English language. Meeting abstracts were excluded. The search strategy is displayed in Supplementary Material 1.

Reference tests

A reference test is the best available test and is the standard against which other assays are compared. Cell cytotoxicity neutralization assay (CCNA) and toxigenic culture (TC) are regarded as reference tests for diagnosing CDI. (2)

CCNA demonstrates the presence of free toxin B. For this test, stool filtrates are inoculated onto a monolayer of a cell culture which is then observed for a toxin B-induced cytopathic effect (rounding of the cells). The cytopathic effect is evaluated at 24 and 48 hours. Cell lines commonly used for CCNA include Vero cells, HeLa cells, human foreskin fibroblast cells and Hep-2 cells. Neutralization of the cytopathic effect is necessary to determine the specificity of this effect and can be done by using *Clostridium sordelli* antitoxin or *C. difficile* antitoxin. (3) This reference test takes 1 to 2 days to perform and requires cell culture and laboratory expertise, so it is not routinely used in most diagnostic laboratories.

TC demonstrates the presence of *C. difficile*, which is able to produce toxins in vitro. Stools are incubated anaerobically for at least 48 hours on selective media. Many different culture media exist for this purpose, all aiming to enhance the recovery of *C. difficile* while inhibiting the overgrowth of other faecal flora. (4) Pretreatment with alcohol shock (5) or heat shock can also be used to decrease overgrowth of normal faecal flora. (4) Also, broth enrichment before plating onto a solid medium is sometimes used (also called enriched culture). (4) Furthermore, a chromogenic medium (ChromID agar; bioMérieux) for the recovery of *C. difficile* has been developed which is designed to isolate and identify *C. difficile* within 24 hours. However, no consensus exists on which culture medium and/or culture method is the most appropriate to use. Colonies suspicious for *C. difficile* can be recognized by Gram staining, colony morphology, 'horse manure' odour, biochemical testing, gas-liquid chromatography, ultraviolet light fluorescence, latex agglutination and matrix-assisted desorption ionization-time of flight mass spectrometry. (6) Isolates from positive cultures are either tested for in vitro toxin production by the use of CCNA or toxin A/B enzyme immunoassay (EIA) or tested for the presence of toxin A/B genes by NAAT.

Index tests

Index tests are the tests whose performance is being evaluated compared to the reference tests. The index tests we reviewed comprise all commonly applied and commercially available laboratory tests for diagnosing CDI other than the reference tests. These include EIAs that detect glutamate dehydrogenase (GDH), EIAs that detect toxins A and B and NAAT.

GDH EIAs detect glutamate dehydrogenase, an enzyme that is produced by both toxigenic and nontoxigenic strains of *C. difficile*. GDH EIAs are available in well-type format (results are displayed as a colour change which can be detected visually or photospectrometrically) or membrane-type format (results can be visually read from a membrane).

Toxin A/B EIAs detect toxins A and B and are also available in well-type or membrane-type format. Most EIAs detecting only toxin A have been replaced by EIAs detecting both toxins A and B, as strains that only produce toxin B and not toxin A are reported.

Several membrane-type tests that include both an EIA detecting GDH and an EIA detecting toxins A and B are also available (*C. diff* Quik Chek Complete, Techlab, Combo *C. difficile*; Theradiag).

NAATs include assays that use PCR, helicase-dependent amplification and loop-mediated isothermal amplification. Most assays detect conserved regions within the gene for toxin B (*tcdB*), but assays that detect a highly conserved sequence of the toxin A gene (*tcdA*) have also been developed (Illumigene, Meridian, Bioscience and Amplivue, Quidel). (7, 8) NAATs that not only detect *tcdB* but also the binary toxin genes (*cdt*) and the deletion at nucleotide 117 on *tcdC* are also available (*Verigene C. difficile* test, Nanosphere and Xpert, Cepheid) and offer the potential advantage of detecting PCR ribotype 027, although highly related PCR ribotypes may also be detected by these tests (without distinguishing them from PCR ribotype 027). (9) NAATs that detect multiple targets at the same time, including *C. difficile* toxin genes, are also available (Seeplex Diarrhea ACE detection, Seegene, xTAG Gastrointestinal Pathogen Panel, Luminex, FilmArray Gastrointestinal Panel, BioFire Diagnostics).

Test performance

The numbers of truly positive, falsely positive, falsely negative and truly negative index test results are generally displayed in a 2 x 2 table (Table 1). Test performance can be derived from this 2 x 2 table. The sensitivity of a test is defined as the probability that the index test result will be positive in a person with disease ($a/a+c$). The specificity of a test is defined as the probability that the index test result will be negative in a person without disease ($d/b+d$). The positive predictive value (PPV) of a test is the probability that a person has the disease, given the positive test result ($a/a+b$). The negative predictive value (NPV) of a test is the probability that a person is free of disease, given the negative test result ($d/c+d$). PPV and NPV are dependent on disease prevalence in the tested population (<http://training-old>).

cochrane.org/sites/training-old.cochrane.org/files/uploads/DTA/1.3_Introduction_to_test_accuracy/story.html).

Table 1. 2x2 table used to calculate test characteristics

	Diseased or reference test positive	Non-diseased or reference test negative
Index test positive	(a) True positive	(b) False positive
Index test negative	(c) False negative	(d) True negative

Eligibility criteria

Studies eligible for inclusion had to: (1) describe original research, (2) compare an index test (one commercially available in Europe) with a reference test (CCNA or TC), (3) perform the tests on *C. difficile*-negative and -positive clinical human stool samples and (4) provide sufficient information to recalculate sensitivity and specificity and their confidence intervals. Culture without determining the toxigenic status was accepted as a reference test if only assays detecting GDH were evaluated.

Studies were excluded if: (1) the reference test was not performed on all samples but only on positive, negative or discordant samples (to exclude partial verification bias), (2) not all samples were tested by the same reference test, (3) the reference method was a composite of more than one test, (4) the reference method included clinical data for its interpretation, (5) the index test was partly used as reference method, (6) the index test did not follow manufacturers' instructions for testing or sample collection, (7) for CCNA, samples were not stored correctly before testing (refrigerated or frozen at -20°C and thawed only once) or neutralization to determine the specificity of the cytopathic effect was not executed and (8) only selected samples were included.

Selection process

Study eligibility was assessed in a two-step selection process by two independent investigators (MC, ET). Inconsistencies were resolved by consensus and by consultation of a third and fourth investigator (EK, TP).

Outcome measures, data extraction and quality assessment

The principal measures of outcome were the sensitivity and specificity of different index tests compared to one of the 2 reference tests. Toxin A/B EIAs, GDH EIAs and NAATs were compared to CCNA and TC. GDH EIAs were additionally compared to culture. From

each study we extracted the number of true-positive, false-positive, false-negative and true-negative findings to be able to calculate the sensitivity and specificity of the index test evaluated in that study. Data were extracted by two independent investigators (MC, ET) using a data extraction form (Supplementary Material 2). Additional data that were extracted included year of publication, storage conditions of the samples, information about the study population and information about the execution of the index test and reference test. The quality of the studies was assessed by the same two independent investigators using a quality assessment tool. This quality assessment tool (Supplementary Material 3) consisted of items from the Quality Assessment for Studies of Diagnostic Accuracy (QUADAS) tool (10), supplemented with items concerning the appropriate handling of specimens and appropriate execution of reference tests.

Statistical analysis

For all index tests in all studies, the sensitivity and specificity and their respective confidence intervals were calculated from the number of true-positive, false-positive, false-negative and true-negative findings supplied in these studies. Wherever possible, the results after initial testing (instead of results after retesting of indeterminate results) were used to calculate the sensitivity and specificity. Random effects logistic regression was used to pool the mean sensitivities and specificities for the different index tests and the different types of index tests. In case of fewer than four studies, a fixed effect model was used. NPVs and PPVs were calculated using a hypothetical prevalence of CDI of 5, 10, 20 and 50% in the tested population. We used Stata 12.0 software (StataCorp) for all statistical analyses.

Guidelines and additional studies

An electronic search was performed on topics concerning laboratory diagnosis of CDI not included in our meta-analysis (e.g. repeated testing, sample selection). Published guidelines on CDI testing were also studied. These included guidelines from the Society for Healthcare Epidemiology of America/Infectious Diseases Society of America (published in 2010) (11), guidelines from the Australasian Society for Infectious Diseases (published in 2011) (12), guidelines from the American College of Gastroenterology (published in 2013) (13), guidelines from the American Academy of Pediatrics (published in 2013) (14) and guidelines from the UK National Health Service (update published in 2012). (15)

Formulation of recommendations

The guideline was developed according to the Appraisal of Guidelines for Research and Evaluation (AGREE II) instrument. (16) Findings of the literature review and meta-analysis results were discussed with the members of the executive committee, and recommendations were formulated. We slightly modified the GRADE system to grade the strength of the recommendations and the quality of evidence (Table 2). (17) A good practice statement could be made instead of a formal graded recommendation for domains where this was deemed appropriate. (18) The drafting group (consisting of experts in the field) and a patients' representative were invited to comment on the recommendations, and results from these discussions were incorporated in the final recommendations.

Table 2. Scoring system for grading quality of evidence and strength of recommendations.

Quality of evidence	
High quality	Evidence from at least 1 properly designed cross sectional or cohort study in patients with diagnostic uncertainty and direct comparison of all test results with an appropriate reference standard.
Moderate quality	Evidence from: (1) at least 1 cross sectional or cohort study in selected patients and/or no or partial comparison of test results with an appropriate reference standard, (2) case control studies
Low quality	Evidence from opinions of respected authorities, based on clinical experience, descriptive case studies or reports of expert committees
Strength of a recommendation	
Strong recommendation for use	Desirable effects clearly outweigh undesirable effects
Weak recommendation for use	Desirable and undesirable effects are closely balanced or recommendation is based on low quality evidence
Weak recommendation against use	Desirable and undesirable effects are closely balanced or recommendation is based on low quality evidence
Strong recommendation against use	Undesirable effects clearly outweigh desirable effects
Good practice statement	Desirable effects clearly outweigh undesirable effects, but no or only indirect evidence is/will become available

Results

Literature search and selection process

A total of 795 unique citations were identified by our current search. On the basis of title and abstract, 693 articles were excluded, leaving 102 full-text articles for detailed assessment. In total, 61 studies were excluded after detailed assessment. Reasons for exclusion were (some studies had more than one reason for exclusion): not all samples were tested by the (same) reference method (23 studies), no or an inadequate reference test was used (16 studies), samples were selected inadequately (13 studies), not enough information was provided (seven studies), the study did not describe original research (five studies), no clinical human stool samples were included (three studies), no commercial diagnostic test was investigated (two studies) and stool samples were incorrectly collected in transport medium (one study).

From all 43 studies included in the previous meta-analysis (1), 28 were excluded. Twenty-four of these studies evaluated tests that were no longer available (mainly EIAs detecting toxin A only). Two other studies were excluded because they did not evaluate a commercial test (both studies evaluated an in-house PCR), one study was excluded because not all samples were tested by the same reference test and one study was excluded because samples were stored incorrectly for CCNA testing. A total of 56 studies (15 from the previous meta-analysis and 41 published since 2009) were included in the meta-analysis. (7, 8, 19-72) A summary of the selection process is shown in Fig. 1.

Study characteristics

Twenty-four different laboratory assays were evaluated: one well-type EIA for GDH, three membrane-type EIAs for GDH, five well-type EIAs for toxins A and B, four membrane-type EIAs for toxin A and B and 11 NAATs (Table 3). In total, 133 comparisons between index tests and reference tests were available, including 53 comparisons to CCNA, 69 comparisons to TC and 11 comparisons to culture. Studies were published between 1996 and 2014. The number of evaluated index tests per study ranged from one to ten, and the number of included samples ranged from 60 to 12 369. The CDI prevalence in the tested population ranged from 6 to 48%. Table 4 lists the characteristics of included studies.

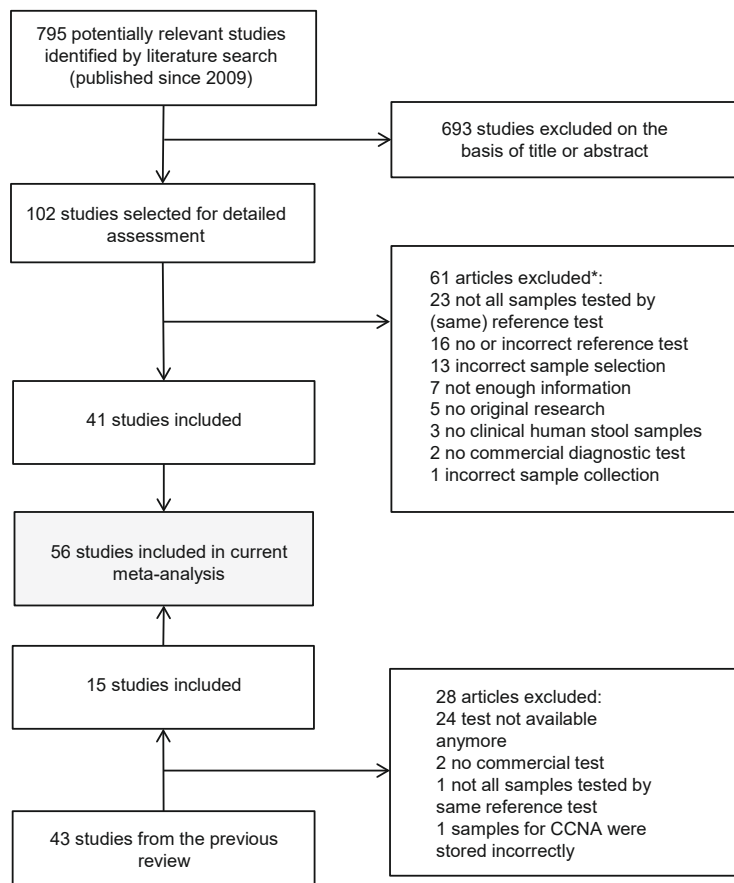


Figure 1. Summary of selection process.

*Some studies had more than one reason for exclusion.

Table 3. Index tests included in meta-analysis

Assay type	Test	Manufacturer	Target(s)	Method
(A) Well-type EIA GDH	C. diff Chek-60	Techlab	GDH	Well-type EIA
(B) Membrane-type EIA GDH	C. diff Quik Chek	Techlab	GDH	Membrane-type EIA
	Immunocard C. <i>difficile</i>	Meridian	GDH	Membrane-type EIA
	Quik Chek Complete-GDH ^a	Techlab	GDH	Membrane-type EIA

Table 3. Continued.

Assay type	Test	Manufacturer	Target(s)	Method
(C) Well-type EIA toxins A&B	Premier tox A/B	Meridian	Toxins A and B	Well-type EIA
	Remel ProSpecT	Oxoid	Toxins A and B	Well-type EIA
	Ridascreen tox A/B	Biopharm	Toxins A and B	Well-type EIA
	<i>Clostridium difficile</i> Tox A/B II	Techlab	Toxins A and B	Well-type EIA
	Vidas CDAB	Biomérieux	Toxins A and B	Automated EIA
(D) Membrane-type EIA toxins A&B	Immunocard tox A/B	Meridian	Toxins A and B	Membrane-type EIA
	Quik Chek Complete-tox A/Ba	Techlab	Toxins A and B	Membrane-type EIA
	Tox A/B Quik Chek	Techlab	Toxins A and B	Membrane-type EIA
	Xpect	Oxoid	Toxins A and B	Membrane-type EIA
(E) NAAT	Advansure CD	LG Life Sciences	<i>tcdA, tcdB</i>	RT-PCR
	Amplivue	Quidel	<i>tcdA</i>	Isothermal helicase-dependent amplification
	BD GeneOhm	Becton, Dickinson	<i>tcdB</i>	RT-PCR
	BD Max Cdiff	Becton, Dickinson	<i>tcdB</i>	RT-PCR
	GenomEra	Abacus Diagnostics	<i>tcdB</i>	RT-PCR
	Illumigene Portrait	Meridian Great Basin	<i>tcdA, tcdB</i>	LAMP
	Prodesse ProGastro Cd Assay	Hologic Gen-Probe	<i>tcdB</i>	Isothermal helicase-dependent amplification
	Seeplex Diarrhea ACE Detection ^c	Seegene	<i>tcdB</i>	RT-PCR
	Verigene	Nanosphere	<i>tcdA, tcdB, cdt^b, tcdC</i> deletion nt 117 ^b	PCR/nanoparticle-based microarray
	Xpert <i>C. difficile</i>	Cepheid	<i>tcdB, cdt, tcdC</i> deletion nt 117	RT-PCR

EIA, enzyme immunoassay; GDH, glutamate dehydrogenase; LAMP, loop-mediated isothermal DNA amplification; RT-PCR, real-time PCR.

^a Part of an EIA that detects both toxins A/B and GDH

^b Only for epidemiologic purposes

^c Multiplex PCR system

Table 4. Characteristics of included studies

Author	Year	Country	Reference test	Index test(s)	Total no. samples	Study population	Consistency stool samples	Prev. CDI (CCNA)	Prev. CDI (TC)
Barikin	2012	US	TC	Premier tox A/B, Immunocard <i>C. difficile</i> , Illumigene	272	Adult inpatients of a large community teaching hospital with diarrhea, risk factors for CDI and for whom a CDI test was requested by their physician	unformed	13.1	
Berg, van den	2005	The Netherlands	CCNA	Immunocard tox A/B	367	unformed stools of adults with a specific request for CDI testing or hospitalized >72hrs that were submitted to the laboratories of 3 university hospitals	unformed	6.3	
Berg, van den	2007	The Netherlands	CCNA	Premier tox A/B	540	unformed stools of patients suspected of having CDI or hospitalized >72hrs in 4 university medical centres	unformed	5.7	
Berry	2014	UK	CCNA	Xpert	1034	inpatients in two acute care hospitals > 15yrs old with suspected CDI for whom CDI testing was requested by the treating physician	unformed	6.0	
Boer, de	2010	The Netherlands	CCNA	Xpect	161	clinical stool specimens from patients for whom a request for CDI testing was issued, prospectively collected at a laboratory for infectious diseases	unclear	9.9	

Table 4. Continued.

Author	Year	Country	Reference test	Index test(s)	Total no. samples	Study population	Consistency stool samples	Prev. CDI (CCNA)	Prev. CDI (TC)
Bruins	2012	The Netherlands	TC	Immunocard tox A/B, Quik Chek Complete, Premier tox A/B, Illumigene	986	hospitalized and non-hospitalized patients with diarrhea who had a stool sample sent to the laboratory of a major hospital, preferably from those patients known to have CDI-associated symptoms or risk factors	unformed		7.4
Buchan	2012	US	TC	Portrait, GeneXpert, GeneOhm, Illumigene	540/275/ 169/96	stool specimens from patients > 2yrs old suspected of having CDI collected at 4 institutions	unformed		22.5
Calderaro	2013	Italy	TC	Illumigene, Quik Chek Complete	306	patients attending a university hospital with a suspicion of CDI	unclear		19.6
Carroll	2013	US	TC	Verigene	1875	leftover stool samples submitted specifically for CDI testing according to the institution's routine practice to 5 geographically diverse clinical microbiology laboratories	formed and unformed		8.4 (direct), 14.7 (enriched)

Table 4. Continued.

Author	Year	Country	Reference test	Index test(s)	Total no. samples	Study population	Consistency stool samples	Prev. CDI (CCNA)	Prev. CDI (TC)
Eastwood	2009	UK	CCNA	Premier tox A/B, Xpect, Tox A/B Quik Chek, Ridascree tox A/B, tox A/B II, ProSpect, VIDAS CDAB, Immunocard tox A/B, C. diff Chek-60, BD GeneOhm	488	stool specimens submitted for CCNA testing at the laboratory of a teaching hospital, 10 samples were randomly chosen each day	unformed	18.1	
Eckert	2014	France	CCNA, TC	Amplivue, C. diff Quik Chek	308	inpatients in 4 university-affiliated hospitals >2yrs old with suspected CDI for whom CDI testing was requested by the treating physician or if diarrhoea occurred after day 3 of hospitalization	unformed	7.5	11.7
Fenner	2008	Switzerland	culture	C. diff chek-60	1468	stools of adults patients suspected of having CDI at a university hospital	unclear		12.7 culture positive
Hart	2014	Australia	culture, TC	Illumigene, BD GeneOhm, Quik Chek Complete	150	stools of children collected at the laboratory of a pediatric hospital fulfilling the criteria for CDI testing in this hospital ^a	formed (4%) and unformed (96%)		30.0

Table 4. Continued.

Author	Year	Country	Reference test	Index test(s)	Total no. samples	Study population	Consistency stool samples	Prev. CDI (CCNA)	Prev. CDI (TC)
Hirvonen	2013	Finland	TC	GenomEra	310	stool specimens from inpatients (7-95yrs old), collected prospectively according to routine hospital practice for antibiotic-associated diarrhea at a large teaching hospital	unformed		24.9
Huang	2009	Sweden	CCNA	Xpert	220	consecutive stool specimens from patients >2yrs old and who were symptomatic and had a request for CDI testing at a university hospital	unformed	10.5	
Jacobs	1996	Israel	culture, TC	Immunocard C. <i>difficile</i>	258	stool samples from patients who developed diarrhea during hospitalization in a community teaching hospital and control samples from 24 patients without diarrhea	formed and unformed		7.0
Jong	2012	The Netherlands	TC	Immunocard tox A/B, VIDAS CDAB	150	hospitalized adult patients in a tertiary teaching hospital who had a stool specimens submitted for CDI testing	unclear		9.7
Kawada	2011	Japan	culture, TC	Quik Chek Complete, Immunocard C. <i>difficile</i> , Tox A/B Quik Chek	60	patients hospitalized at a geriatric hospital and diagnosed as having antibiotic-associated diarrhea	unformed		46.7

Table 4. Continued.

Author	Year	Country	Reference test	Index test(s)	Total no. samples	Study population	Consistency stool samples	Prev. CDI (CCNA)	Prev. CDI (TC)
Kim	2014	Korea	TC	Quik Chek Complete, VIDAS CDAB	608	suspected CDI patients in a tertiary care teaching hospital	unformed		9.0
Kim	2012	Korea	TC	AdvanSure, VIDAS CDAB	127	diarrhoeal stool specimens submitted to a hospital laboratory for <i>C. difficile</i> culture	unformed		8.8
Lalande	2011	France	TC	Illumigene	472	consecutive stools from patients suspected of having CDI	unformed		10.4
Larson	2010	US	CCNA	C. diff Quik Chek	699	stool samples submitted for CDI testing from adult patients at a university hospital	unformed	6.7	
Le Guern	2012	France	TC	BD Max Cdiff, BD GeneOhm, Tox A/B Quik Chek	360	diarrheal stool specimens collected from inpatients at a university hospital	unformed		12.2
Leitner	2013	Austria	TC	BD Max Cdiff, Premier tox A/B	180	stool specimens from adults and children with a specified request for CDI testing at a medical university	unformed		16.7
Massey	2003	Canada	CCNA	Tox A/B II	557	stools samples of adult hospitalized patients suspected of having CDI at a large teaching hospital	unformed	25.7	
Mattner	2012	Germany	TC	Ridascreen	254	all liquid stool samples sent to a university microbiology laboratory	unformed		16.4

Table 4. Continued.

Author	Year	Country	Reference test	Index test(s)	Total no. samples	Study population	Consistency stool samples	Prev. CDI (CCNA)	Prev. CDI (TC)
Musher	2007	US	CCNA	Premier tox A/B, Immunocard tox A/B, Tox A/B II, ProSpect	446/131	consecutive stool samples submitted to the laboratory of a medical center for CDI testing	unclear	17.0/41.2	
Noren	2011	Sweden	CCNA	llumigene	272	consecutive stool specimens from adults and children submitted for CDI testing from hospitals and communities	unclear	13.2	
Novak-Weekley	2010	US	TC	Xpert, Premier A/B	432	leftover stool samples from patients >2yrs old with suspected CDI for whom toxin EIAs were ordered according to the institution's standard practices at regional reference laboratories serving hospitals and associated medical clinics	unformed		16.8
O'Connor	2001	Ireland	CCNA	Tox A/B II, Premier tox A/B	200	consecutive stools of adult patients suspected of having CDI submitted to the laboratories of university hospitals	formed and unformed	30.5	
Ota	2012	USA	CCNA	C. diff Quik Chek Complete, Premier tox A/B, Illumigene	141	consecutive stool specimens prospectively collected at a children's hospital from patients 1-18 years of age and submitted for CDI testing	unformed	18.4	

Table 4. Continued.

Author	Year	Country	Reference test	Index test(s)	Total no. samples	Study population	Consistency stool samples	Prev. CDI (CCNA)	Prev. CDI (TC)
Pancholi	2012	USA	CCNA	Illumigene, Xpert	200	consecutive and prospectively collected stools from adult patients submitted to a university medical center laboratory for routine CDI testing.	unformed	11.6	
Planche	2013	UK	CCNA, TC	Xpert, C. diff Chek-60, Premier tox A/B, Tox A/B II	8827/12365/ 9192/12369	faecal samples from hospital and community patients submitted for routine CDI testing according to a routine protocol ¹⁸ submitted to 4 hospital diagnostic laboratories serving major teaching hospitals and their communities	unformed	5.9	8.4
Qutub	2011	Saudi Arabia	CCNA	C. diff Chek-60, Tox A/B II	150	stool samples from consecutive inpatients with suspected CDI	unformed	34.7	
Reller	2007	US	culture	C. diff chek-60	439	stool samples from hospitalized adults and children suspected of having CDI	unclear		36.7 culture positive
Reller	2010	US	CCNA	C. diff Chek-60, Quik Chek, Tox A/B Quik Chek	600	sequential weekday stool samples submitted to a university hospital microbiology laboratory for CDI testing	unformed	7.7	
Shin	2009	Korea	TC	Vidas CDAB	1596	stool samples from patients admitted to a tertiary teaching hospital with clinical signs compatible with CDI			19.6

Table 4. Continued.

Author	Year	Country	Reference test	Index test(s)	Total no. samples	Study population	Consistency stool samples formed (51% and unformed	Prev. CDI (CCNA)	Prev. CDI (TC)
Shin	2009	Korea	TC	Vidas CDAB	555	patients >2yrs old with suspected CDI from two hospitals	formed (51% and unformed	20.3	
Shin	2012	Korea	TC	Seegene, BD GeneOhm	243	fresh stool specimens from patients with clinical signs compatible with CDI who were hospitalized in 3 teaching hospitals	unclear	28.8	
Shin	2012	Korea	TC	Xpert/epi, Vidas CDAB	253	consecutive stool specimens from suspected CDI patients in a tertiary hospital	unformed	18.4	
Sloan	2008	US	TC	Premier tox A/B, Xpect, Immunocard A/B	200	stools of patients suspected of having CDI submitted to a clinical microbiology laboratory of a large tertiary care teaching hospital	unformed	22.0	
Snell	2004	Canada	culture, TC	C. diff chek-60, Tox A/B II	497	stools of inpatients suspected of having CDI at a large teaching hospital	unformed	10.5	
Soh	2014	Korea	TC	AdvanSure CD, Illumigene	203	stool samples collected at a tertiary university teaching hospital	unformed	12.8	
Stamper	2009	US	CCNA	BD GeneOhm	401	symptomatic adults patients who had a stool sample submitted for routine CDI testing in a tertiary care university medical center	unformed	11.0	

Table 4. Continued.

Author	Year	Country	Reference test	Index test(s)	Total no. samples	Study population	Consistency stool samples	Prev. CDI (CCNA)	Prev. CDI (TC)
Stamper	2009	US	CCNA, TC	ProGastro CD	280	stool samples submitted for routine CDI testing from symptomatic patients >2yrs old at a tertiary-care university medical institution	unformed	11.0	15.7
Staneck	1996	US	CCNA	Immunocard C. <i>difficile</i>	906	stool samples submitted to 3 hospital microbiology laboratories	unclear	14.1	
Swindells	2010	UK	culture, CCNA, TC	C. diff Quik Chek Complete, Vidas CDAB, Xpert, GeneOhm	150	consecutive stool specimens from inpatients >65 yrs old who developed diarrhea at least 48hrs after admission	unformed	10.0	12.0
Tenover	2010	US/Canada	TC	Xpert	2296	leftover stool specimens from patients >2yrs old from 7 health care organizations (6 US, 1 Canada) for whom CDI testing was ordered according to the institution's practices	unformed		10.8 (direct), 14.7 (enriched)
Terhes	2009	Hungary	CCNA	BD GeneOhm	600	inpatients and outpatients at a local university hospital who had a diarrheal stool sample sent to the laboratory for CDI testing	unformed	6.4	

Table 4. Continued.

Author	Year	Country	Reference test	Index test(s)	Total no. samples	Study population	Consistency stool samples	Prev. CDI (CCNA)	Prev. CDI (TC)
Ticehurst	2006	US	CCNA	C. diff chek-60	266	stools of patients suspected of having CDI submitted to the laboratories of two acute-care hospitals	unclear	9.0	
Turgeon	2003	US	CCNA	Immunocard C. difficile	1003	consecutive stools of adults and children suspected of having CDI at 5 major hospitals	unformed and formed	10.1	
Vanpoucke	2001	Belgium	CCNA	Ridascreen	156	stool specimens submitted to the laboratory of a university hospital with a request for CDI testing	unformed	31.8	
Viala	2012	France	TC	BD GeneOhm, Xpert, Illumigene	94	fresh stool specimens from symptomatic patients collected at a university hospital, 45 TC+ and 49 TC- were selected	unformed		47.8
Walkty	2013	Canada	TC	Illumigene, C. diff Quik Chek	428	all diarrheal stool specimens from patients >1yrs old submitted for CDI testing to 3 microbiological laboratories serving major hospitals and surrounding communities	unformed		14.7
Wren	2009	UK	culture, TC	C. diff Quik Chek, Tox A/B Quik Chek	1007	stool samples submitted for CDI testing from patients who developed diarrhoea after being admitted to major university hospitals	unformed		8.6

Table 4. Continued.

Author	Year	Country	Reference test	Index test(s)	Total no. samples	Study population	Consistency stool samples	Prev. CDI (CCNA)	Prev. CDI (TC)
Zheng	2004	US	culture	C. diff chek-60	992	stool samples submitted for routine CDI testing because of antibiotic-associated diarrhea collected from hospital laboratories and supplied to TechLab, a large medical center and the reference laboratory	unclear	13.8	

CCNA, cell cytotoxicity neutralization assay; CDI, *Clostridium difficile* infection; TC, toxigenic culture

^aCriteria were: oncology/haematology patient, specific request for CDI testing by treating physician, history of diarrhoea developed while receiving antibiotics, or pseudomembranous colitis

^bCriteria were: all unformed faecal samples not clearly attributable to an underlying disease, or treatment from all hospital patients >2years and from individuals in the community >65 years irrespective of *C. difficile* or other testing requests

Quality assessment

None of the studies fulfilled all our quality assessment criteria, mainly because required information was frequently missing (Fig. 2, Supplementary Material 4). The process used to select samples was adequately reported in 23 (41%) of 56 studies. A minority of studies (6/56, 11%) reported that they did not exclude formed samples from CDI testing. In around half of the studies, conditions of storage for the samples before testing with the index test were not (or were insufficiently) reported. Samples tested by GDH EIA, toxin A/B EIA and NAAT were reported to be stored according to manufacturer's instructions in 10 (46%) of 22, 14 (45%) of 31 and 15 (50%) of 30 studies, respectively. In the remaining 12, 16 and 15 studies, respectively, storage conditions did not or not completely comply with manufacturer's instructions. In 18 (72%) of 25 studies using CCNA as the reference test, samples were stored according to our predefined storage requirements: samples were either refrigerated and tested within 5 days (15 studies) (8, 25, 27, 36, 45-48, 58-61, 63, 65, 68) or were frozen at -20°C and thawed no more than once (three studies) (44, 66, 67). In the remaining seven studies (28%), storage conditions for CCNA were not or incompletely described. Storage conditions for samples tested by TC were reported in 23 (68%) of 34 studies, but no specific requirements for storage of samples tested by TC were set. The execution of the reference test was described in sufficient detail in 44 (79%) of 56 studies. In 2 (8%) of 26 studies using CCNA as reference test, the incubation period was only 24 hours. (61, 63) In studies using TC as reference test, ethanol shock was reported to be performed in 18 of 35 studies (19, 21, 23, 32, 35, 37, 38, 47, 51-55, 57, 61, 69-71), and heat shock was performed in three of 35 studies. (22, 49, 58) Eight studies (23%) used an enrichment broth before plating onto a solid agar. (19, 22-24, 32, 43, 58, 62) Toxigenicity was confirmed by PCR (15/32, 47%) (21, 23, 29, 33-35, 37, 51-57, 70), CCNA (9/32, 28%) (7, 8, 22, 24, 43, 47, 58, 61, 62), toxin EIA (7/32, 22%) (19, 30, 32, 38, 40, 69, 71) or both PCR and CCNA (1/32, 3%). (26) Blinding (index test interpreted without knowledge of reference test or vice versa) was reported in 8 (14%) of 56 studies. Thirty-one studies (55%) reported if any indeterminate results (i.e. invalid, 'no call' or difficult-to-interpret results) were found. Indeterminate results actually occurred in 28 studies and were reported for one membrane-type GDH EIA (ImmunoCard C. difficile), three membrane-type toxin A/B EIAs (Tox A/B Quik Chek, ImmunoCard Tox A/B, Xpect), one automated EIA (Vidas) and nine NAATs. The amount of indeterminate results ranged from 0.3 to 6.8% of tested samples. Repeat testing of samples after an initial indeterminate result was done in 24 (86%) of these 28 studies. Of these, 22 presented results only after repeat testing (7, 8, 20, 21, 24, 29, 30, 34, 35, 37, 38, 43, 46, 47, 54, 58, 59, 62, 65, 69, 70), and two presented results of both initial and repeat testing. (27, 63)

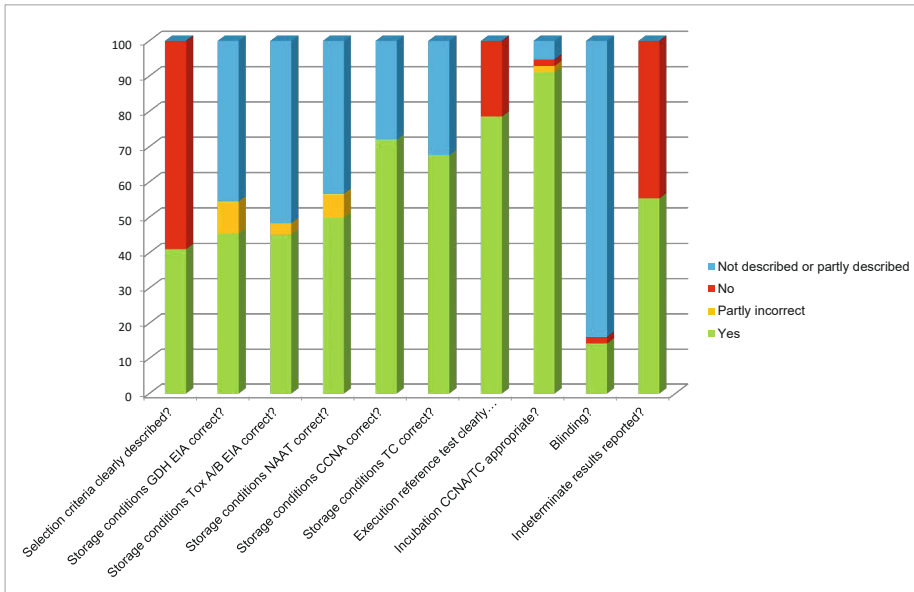


Figure 2. Quality assessment of included studies.

Test performances

Sensitivity and specificity of the index tests were calculated on the basis of the numbers provided in the articles. Discrepancies between calculated sensitivity or specificity and published data were found in two articles; the correct data were provided by both authors upon request. (38, 39) In Table 5, sensitivity and specificity of index tests are compared to CCNA. Reported estimates of sensitivity ranged from 0.80 to 1.00 for GDH EIAs, from 0.44 to 0.99 for toxin A/B EIAs and from 0.83 to 1.00 for NAATs. Reported estimates of specificity ranged from 0.82 to 0.95 for GDH EIAs, from 0.87 to 1.00 for toxin A/B EIAs and from 0.87 to 0.98 for NAATs. Table 6 lists sensitivity and specificity compared to TC. Sensitivities ranged from 0.83 to 1.00, 0.29 to 0.86 and 0.77 to 1.0 for GDH EIAs, toxin A/B EIAs and NAATs, respectively. Specificities ranged from 0.88 to 1.00, 0.91 to 1.00 and 0.83 to 1.00, respectively. In Table 7, sensitivity and specificity of GDH EIAs are compared to culture. Sensitivities ranged from 0.71 to 1.00, and specificities ranged from 0.67 to 1.00. In Table 8, estimates of pooled sensitivity and pooled specificity for the different categories of index tests are shown. The estimated pooled sensitivities and specificities compared to CCNA were used to compute PPVs and NPVs of the categories of index tests at different hypothetical CDI prevalences (Table 9, Supplementary Material 5). At a CDI prevalence of

5%, PPVs ranged from 34 to 81%, and NPVs ranged from 99 to 100%. At a CDI prevalence of 50%, PPVs ranged from 91 to 99%, while NPVs ranged from 83 to 98%.

Table 5. Sensitivity and specificity of index tests compared to CCNA

Index test	Study	Sensitivity (95% CI)	Specificity (95% CI)
(A) Well-type EIA GDH			
<i>C. diff</i> Chek-60	Eastwood [27]	0.90 (0.82–0.95)	0.93 (0.90–0.95)
<i>C. diff</i> Chek-60	Planche [47]	0.96 (0.95–0.98)	0.92 (0.92–0.93)
<i>C. diff</i> Chek-60	Qutub [48]	0.94 (0.84–0.99)	0.88 (0.80–0.94)
<i>C. diff</i> Chek-60	Reller [50]	0.91 (0.79–0.98)	0.90 (0.87–0.92)
<i>C. diff</i> Chek-60	Ticehurst [64]	0.96 (0.79–1.00)	0.90 (0.86–0.94)
(B) Membrane-type EIA GDH			
<i>C. diff</i> Quik Chek	Eckert [8]	1.00 (0.85–1.00)	0.92 (0.88–0.94)
<i>C. diff</i> Quik Chek	Larson [36]	1.00 (0.92–1.00)	0.90 (0.87–0.92)
<i>C. diff</i> Quik Chek	Reller [50]	1.00 (0.92–1.00)	0.83 (0.79–0.86)
ImmunoCard <i>C. difficile</i>	Staneck [60]	0.84 (0.77–0.90)	0.92 (0.90–0.94)
ImmunoCard <i>C. difficile</i>	Turgeon [65]	0.80 (0.71–0.87)	0.92 (0.91–0.94)
Quik Chek Complete-GDH	Ota [45]	0.81 (0.61–0.93)	0.82 (0.73–0.88)
Quik Chek Complete-GDH	Swindells [61]	1.00 (0.78–1.00)	0.95 (0.90–0.98)
(C) Well-type EIA toxins A/B			
<i>Clostridium difficile</i> Tox A/B II	Eastwood [27]	0.91 (0.84–0.95)	0.96 (0.93–0.97)
<i>Clostridium difficile</i> Tox A/B II	Massey [39]	0.75 (0.67–0.82)	0.98 (0.96–0.99)
<i>Clostridium difficile</i> Tox A/B II	Musher [41]	0.96 (0.87–1.00)	0.87 (0.77–0.94)
<i>Clostridium difficile</i> Tox A/B II	O'Connor [44]	0.80 (0.68–0.89)	0.99 (0.96–1.00)
<i>Clostridium difficile</i> Tox A/B II	Planche [47]	0.83 (0.80–0.86)	0.99 (0.99–0.99)
<i>Clostridium difficile</i> Tox A/B II	Qutub [48]	0.73 (0.59–0.84)	1.00 (0.96–1.00)
Premier toxins A/B	Berg, van den 2007 [67]	0.97 (0.83–1.00)	0.94 (0.92–0.96)
Premier toxins A/B	Eastwood [27]	0.92 (0.85–0.96)	0.97 (0.95–0.98)
Premier toxins A/B	Musher [41]	0.99 (0.93–1.00)	0.97 (0.95–0.99)
Premier toxins A/B	O'Connor [44]	0.82 (0.70–0.91)	0.99 (0.96–1.00)
Premier toxins A/B	Ota [45]	0.58 (0.37–0.77)	1.00 (0.97–1.00)
Premier toxins A/B	Planche [47]	0.67 (0.63–0.71)	0.99 (0.99–0.99)
Remel ProSpecT	Eastwood [27]	0.90 (0.83–0.95)	0.93 (0.90–0.95)
Remel ProSpecT	Musher [41]	0.91 (0.80–0.97)	0.97 (0.91–1.00)
Ridascreen toxins A/B	Eastwood [27]	0.67 (0.57–0.75)	0.95 (0.93–0.97)
Ridascreen toxins A/B	Vanpoucke [68]	0.57 (0.43–0.70)	0.97 (0.92–0.99)
(D) Membrane-type EIA toxins A/B			
ImmunoCard toxins A/B	Berg, van den (2005) [66]	0.91 (0.72–0.99)	0.97 (0.95–0.99)
ImmunoCard toxins A/B	Eastwood [27]	0.85 (0.76–0.91)	0.99 (0.98–1.00)
ImmunoCard toxins A/B	Musher [41]	0.96 (0.89–0.99)	0.99 (0.97–1.00)
Quik Chek Complete-Tox A/B	Ota [45]	0.50 (0.30–0.70)	1.00 (0.97–1.00)
Quik Chek Complete-Tox A/B	Swindells [61]	0.73 (0.45–0.92)	1.00 (0.97–1.00)

Table 5. Continued.

Index test	Study	Sensitivity (95% CI)	Specificity (95% CI)
Tox A/B Quik Chek	Eastwood [27]	0.84 (0.76–0.91)	0.99 (0.98–1.00)
Tox A/B Quik Chek	Reller [50]	0.61 (0.45–0.75)	0.99 (0.98–1.00)
Xpect	Boer, de [25]	0.44 (0.20–0.70)	1.00 (0.97–1.00)
Xpect	Eastwood [27]	0.83 (0.74–0.90)	0.99 (0.98–1.00)
(E) Automated EIA toxins A/B			
VIDAS CDAB	Eastwood [27]	0.98 (0.93–1.00)	0.99 (0.98–1.00)
VIDAS CDAB	Swindells [61]	0.53 (0.27–0.79)	1.00 (0.97–1.00)
(F) NAAT			
Amplivue	Eckert [8]	0.96 (0.78–1.00)	0.95 (0.91–0.97)
BD GeneOhm	Eastwood [27]	0.92 (0.85–0.97)	0.95 (0.93–0.97)
BD GeneOhm	Stamper (2009–1) [59]	0.91 (0.78–0.97)	0.95 (0.92–0.97)
BD GeneOhm	Swindells [61]	1.00 (0.78–1.00)	0.98 (0.94–1.00)
BD GeneOhm	Terhes [63]	0.95 (0.82–0.99)	0.96 (0.94–0.98)
Illumigene	Noren [42]	1.00 (0.90–1.00)	0.93 (0.89–0.96)
Illumigene	Ota [45]	0.88 (0.70–0.98)	0.97 (0.93–0.99)
Illumigene	Pancholi [46]	0.87 (0.66–0.97)	0.91 (0.86–0.95)
Prodesse ProGastro Cd assay	Stamper (2009–2) [58]	0.83 (0.65–0.94)	0.96 (0.92–0.98)
Xpert <i>C. difficile</i>	Berry [20]	1.00 (0.94–1.00)	0.94 (0.92–0.95)
Xpert <i>C. difficile</i>	Huang [31]	0.96 (0.78–1.00)	0.87 (0.82–0.92)
Xpert <i>C. difficile</i>	Pancholi [46]	1.00 (0.85–1.00)	0.89 (0.83–0.93)
Xpert <i>C. difficile</i>	Planche [47]	0.98 (0.96–0.99)	0.93 (0.92–0.94)
Xpert <i>C. difficile</i>	Swindells [61]	1.00 (0.78–1.00)	0.97 (0.93–0.99)

CI, confidence interval; CCNA, cell cytotoxicity neutralization assay; EIA, enzyme immunoassay; GDH, glutamate dehydrogenase; NAAT, nucleic acid amplification test

Table 6. Sensitivity and specificity of index tests compared to TC

Index test	Study	Sensitivity (95% CI)	Specificity (95% CI)
(A) Well-type EIA GDH			
<i>C. diff</i> Chek-60	Planche [47]	0.94 (0.93–0.96)	0.94 (0.94–0.95)
(B) Membrane-type EIA GDH			
<i>C. diff</i> Quik Chek	Eckert [8]	0.97 (0.85–1.00)	0.95 (0.92–0.97)
<i>C. diff</i> Quik Chek	Walkty [70]	0.83 (0.71–0.91)	0.97 (0.95–0.98)
ImmunoCard <i>C. difficile</i>	Barkin [19]	1.00 (0.90–1.00)	1.00 (0.98–1.00)
ImmunoCard <i>C. difficile</i>	Jacobs [32]	0.60 (0.32–0.84)	0.76 (0.68–0.83)
Quik Chek Complete—GDH	Bruins [21]	0.97 (0.90–1.00)	0.98 (0.96–0.98)
Quik Chek Complete—GDH	Kawada [33]	1.00 (0.88–1.00)	0.88 (0.71–0.96)
Quik Chek Complete—GDH	Swindells [61]	1.00 (0.81–1.00)	0.97 (0.92–0.99)
(C) Well-type EIA toxins A/B			
<i>Clostridium difficile</i> Tox A/B II	Planche [47]	0.58 (0.55–0.61)	0.99 (0.98–0.99)
<i>Clostridium difficile</i> Tox A/B II	Snell [56]	0.85 (0.72–0.93)	0.98 (0.96–0.99)
Premier toxins A/B	Barkin [19]	0.86 (0.71–0.95)	0.91 (0.86–0.94)
Premier toxins A/B	Bruins [21]	0.41 (0.30–0.53)	0.99 (0.98–0.99)
Premier toxins A/B	Leitner [38]	0.40 (0.21–0.61)	1.00 (0.98–1.00)
Premier toxins A/B	Novak-Weekley [43]	0.58 (0.46–0.70)	0.95 (0.92–0.97)
Premier toxins A/B	Planche [47]	0.46 (0.42–0.49)	0.99 (0.99–0.99)
Premier toxins A/B	Sloan [55]	0.48 (0.32–0.63)	0.98 (0.94–1.00)
Ridascreen toxins A/B	Mattner [40]	0.52 (0.36–0.68)	0.98 (0.95–0.99)
(D) Membrane-type EIA toxins A/B			
ImmunoCard toxins A/B	Bruins [21]	0.41 (0.30–0.53)	0.99 (0.98–1.00)
ImmunoCard toxins A/B	de Jong [26]	0.47 (0.23–0.72)	0.99 (0.96–1.00)
ImmunoCard toxins A/B	Sloan [55]	0.48 (0.32–0.63)	0.99 (0.95–1.00)
Quik Chek Complete—Tox A/B	Bruins [21]	0.55 (0.43–0.66)	1.00 (1.00–1.00)
Quik Chek Complete—Tox A/B	Calderaro [23]	0.68 (0.55–0.80)	0.89 (0.84–0.92)
Quik Chek Complete—Tox A/B	Hart [29]	0.29 (0.16–0.44)	1.00 (0.97–1.00)
Quik Chek Complete—Tox A/B	Kawada [33]	0.79 (0.59–0.92)	0.97 (0.84–1.00)
Quik Chek Complete—Tox A/B	Kim (2014) [35]	0.64 (0.50–0.76)	0.98 (0.96–0.99)

Table 6. Continued.

Index test	Study	Sensitivity (95% CI)	Specificity (95% CI)
Quik Chek Complete— Tox A/B	Swindells [61]	0.61 (0.36–0.83)	1.00 (0.97–1.00)
Tox A/B Quik Chek	Kawada [33]	0.71 (0.51–0.87)	0.94 (0.79–0.99)
Tox A/B Quik Chek	Le Guern [37]	0.43 (0.28–0.59)	1.00 (0.98–1.00)
Tox A/B Quik Chek	Wren [71]	0.40 (0.30–0.51)	1.00 (1.00–1.00)
Xpect	Sloan [55]	0.48 (0.32–0.63)	0.84 (0.77–0.89)
(E) Automated EIA toxins A/B			
VIDAS CDAB	Jong, de [26]	0.71 (0.42–0.92)	0.95 (0.90–0.98)
VIDAS CDAB	Kim (2012) [34]	0.64 (0.31–0.89)	1.00 (0.97–1.00)
VIDAS CDAB	Kim (2014) [35]	0.76 (0.61–0.87)	0.97 (0.96–0.99)
VIDAS CDAB	Shin (2009–1) [52]	0.68 (0.62–0.73)	0.96 (0.95–0.97)
VIDAS CDAB	Shin (2009–2) [51]	0.69 (0.59–0.78)	0.97 (0.94–0.98)
VIDAS CDAB	Shin (2012–2) [54]	0.44 (0.30–0.60)	1.00 (0.98–1.00)
VIDAS CDAB	Swindells [61]	0.44 (0.22–0.69)	1.00 (0.97–1.00)
(F) NAAT			
Advansure CD	Kim (2012) [34]	1.00 (0.72–1.00)	0.98 (0.94–1.00)
Advansure CD	Soh [57]	0.85 (0.65–0.96)	0.98 (0.95–1.00)
Amplivue	Eckert [8]	0.86 (0.71–0.95)	0.98 (0.95–0.99)
BD GeneOhm	Buchan [22]	0.97 (0.86–1.00)	0.98 (0.95–1.00)
BD GeneOhm	Hart [29]	0.89 (0.76–0.96)	0.99 (0.95–1.00)
BD GeneOhm	Le Guern [37]	0.95 (0.85–0.99)	1.00 (0.98–1.00)
BD GeneOhm	Shin (2012–1) [53]	0.96 (0.88–0.99)	0.97 (0.93–0.99)
BD GeneOhm	Swindells [61]	0.94 (0.73–1.00)	0.99 (0.96–1.00)
BD GeneOhm	Viala [69]	0.96 (0.85–0.99)	0.98 (0.89–1.00)
BD Max Cdiff	Le Guern [37]	0.98 (0.88–1.00)	1.00 (0.98–1.00)
BD Max Cdiff	Leitner [38]	0.96 (0.80–1.00)	0.99 (0.96–1.00)
GenomEra	Hirvonen [30]	1.00 (0.95–1.00)	0.99 (0.96–1.00)
llumigene	Barkin [19]	1.00 (0.90–1.00)	1.00 (0.98–1.00)
llumigene	Bruins [21]	0.93 (0.85–0.98)	1.00 (0.99–1.00)
llumigene	Buchan [22]	0.93 (0.68–1.00)	0.95 (0.88–0.99)
llumigene	Calderaro [23]	1.00 (0.94–1.00)	0.83 (0.78–0.87)
llumigene	Hart [29]	0.89 (0.76–0.96)	1.00 (0.97–1.00)
llumigene	Lalande [7]	0.92 (0.80–0.98)	0.99 (0.98–1.00)
llumigene	Soh [57]	0.92 (0.75–0.99)	0.99 (0.97–1.00)
llumigene	Viala [69]	0.87 (0.73–0.95)	1.00 (0.93–1.00)
llumigene	Walkty [70]	0.73 (0.60–0.83)	1.00 (0.98–1.00)
Portrait	Buchan [22]	0.98 (0.94–1.00)	0.93 (0.90–0.95)

Table 6. Continued.

Index test	Study	Sensitivity (95% CI)	Specificity (95% CI)
Prodesse ProGastro Cd assay	Stamper (2009–2) [58]	0.77 (0.62–0.89)	0.99 (0.97–1.00)
Seeplex ACE	Shin (2012–1) [53]	0.90 (0.80–0.96)	0.97 (0.93–0.99)
Verigene	Caroll [24]	0.91 (0.87–0.94)	0.93 (0.91–0.94)
Xpert <i>C. difficile</i>	Buchan [22]	1.00 (0.94–1.00)	0.92 (0.87–0.95)
Xpert <i>C. difficile</i>	Novak-Weekley [43]	0.94 (0.86–0.98)	0.96 (0.94–0.98)
Xpert <i>C. difficile</i>	Planche [47]	0.95 (0.93–0.96)	0.96 (0.96–0.97)
Xpert <i>C. difficile</i>	Shin (2012–2) [54]	1.00 (0.93–1.00)	0.95 (0.91–0.98)
Xpert <i>C. difficile</i>	Swindells [61]	1.00 (0.81–1.00)	0.99 (0.96–1.00)
Xpert <i>C. difficile</i>	Tenover [62]	0.93 (0.90–0.96)	0.94 (0.93–0.95)
Xpert <i>C. difficile</i>	Viala [69]	0.98 (0.88–1.00)	0.98 (0.89–1.00)

CI, confidence interval; EIA, enzyme immunoassay; GDH, glutamate dehydrogenase; NAAT, nucleic acid amplification test; TC, toxigenic culture.

Table 7. Sensitivity and specificity of index tests compared to culture

Index test	Study	Sensitivity (95% CI)	Specificity (95% CI)
(A) Well-type EIA GDH			
<i>C. diff</i> Chek-60	Fenner [28]	0.93 (0.88–0.97)	0.97 (0.95–0.97)
<i>C. diff</i> Chek-60	Reller (2007) [49]	1.00 (0.98–1.00)	0.67 (0.61–0.72)
<i>C. diff</i> Chek-60	Snell [56]	0.94 (0.86–0.98)	0.98 (0.96–0.99)
<i>C. diff</i> Chek-60	Zheng [72]	0.71 (0.63–0.78)	0.88 (0.85–0.90)
(B) Membrane-type EIA GDH			
<i>C. diff</i> Quik Chek	Wren [71]	0.95 (0.90–0.98)	0.99 (0.98–1.00)
Quik Chek Complete—GDH	Bruins [21]	0.95 (0.89–0.99)	0.99 (0.98–0.99)
Quik Chek Complete—GDH	Hart [29]	0.87 (0.75–0.95)	0.97 (0.91–0.99)
Quik Chek Complete—GDH	Kawada [33]	1.00 (0.88–1.00)	0.93 (0.78–0.99)
Quik Chek Complete—GDH	Swindells [61]	1.00 (0.82–1.00)	0.98 (0.93–1.00)
ImmunoCard <i>C. difficile</i>	Jacobs [32]	0.75 (0.59–0.87)	0.90 (0.83–0.95)
ImmunoCard <i>C. difficile</i>	Kawada [33]	0.80 (0.61–0.92)	1.00 (0.88–1.00)

CI, confidence interval; EIA, enzyme immunoassay; GDH, glutamate dehydrogenase; NAAT, nucleic acid amplification test.

Discussion

In the present meta-analysis, we evaluated the diagnostic accuracy of various commercial laboratory assays for diagnosing CDI. Toxin A/B EIAs tended to be the most specific assays, while GDH EIAs and NAATs were more sensitive tests. Although many toxin A/B EIAs belong to the least sensitive tests, the sensitivity of this category of assays is not as low as reported earlier. (1) This is because only currently available tests were included in the present analysis, and the newer generation of toxin A/B EIAs turns out to be more sensitive than the earlier toxin A EIAs.

We compared all categories of the index tests (GDH EIAs, toxin A/B EIAs and NAATs) to both of the reference tests, CCNA and TC. However, not only are the targets of these three categories of index tests somewhat different, but also the targets of the two reference tests differ: CCNA detects in vivo toxin production, while TC detects the presence of a toxigenic *C. difficile* strain.

This explains why sensitivities and specificities were different for each reference test that was used as a comparator. For example, toxin A/B EIAs were less sensitive compared to TC instead of CCNA: toxin EIAs will not (like the TC) detect all samples containing toxigenic *C. difficile* strains but only (some of) those with free toxin present. It also explains why NAATs were less specific compared to CCNA instead of TC: NAATs are not able (like CCNA) to discern samples with in vivo toxin production from samples with in vitro toxin production.

We included both CCNA and TC as reference tests, as there has always been debate which of these tests best defines CDI cases. Recently a large study reported that CCNA positivity (i.e. demonstration of free toxin) but not TC positivity (i.e. demonstration of toxin-producing capacity) correlated with clinical outcome. Therefore, at least all samples with a positive CCNA can be considered to represent true CDI cases. (47) However, samples with a positive TC but negative CCNA are difficult to interpret. These samples could either belong to *C. difficile* carriers (harbouring a toxigenic *C. difficile* strain not producing detectable toxins at that moment) or to patients with CDI with toxin levels below the threshold of detection.

To guarantee a certain level of uniformity and quality, only studies that met our eligibility criteria were included in the meta-analysis. Still, studies differed from one another in many aspects. For CCNA, diverse dilutions of faecal filtrate and diverse cell lines were used. For TC, diverse culture media and diverse methods to demonstrate toxigenicity were applied.

Table 8. Pooled sensitivities and specificities of categories of tests

Type	Test	Compared to CCNA			Compared to TC			Compared to culture		
		No. of studies	Sensitivity (95% CI)	Specificity (95% CI)	No. of studies	Sensitivity (95% CI)	Specificity (95% CI)	No. of studies	Sensitivity (95% CI)	Specificity (95% CI)
EIA GDH	Total	12	0.94 (0.89–0.97)	0.90 (0.88–0.92)	8	0.96 (0.86–0.99)	0.96 (0.91–0.98)	11	0.94 (0.86–0.97)	0.96 (0.92–0.98)
	Well type	5	0.94 (0.91–0.97)	0.92 (0.92–0.93)	1	0.94 (0.93–0.96)	0.94 (0.94–0.95)	4	0.89 (0.86–0.91)	0.91 (0.90–0.92)
	Membrane type	7	0.98 (0.78–1.00)	0.90 (0.87–0.93)	7	0.97 (0.84–1.00)	0.96 (0.90–0.99)	7	0.93 (0.84–0.97)	0.98 (0.95–0.99)
EIA toxins A/B	Total	27	0.83 (0.76–0.88)	0.99 (0.98–0.99)	29	0.57 (0.51–0.63)	0.99 (0.98–0.99)			
	Well type	18	0.85 (0.77–0.91)	0.98 (0.96–0.99)	16	0.60 (0.52–0.68)	0.98 (0.97–0.99)			
	Membrane type	9	0.79 (0.66–0.88)	0.99 (0.98–0.99)	13	0.53 (0.45–0.61)	0.99 (0.97–1.00)			
NAAT		14	0.96 (0.93–0.98)	0.94 (0.93–0.95)	32	0.95 (0.92–0.97)	0.98 (0.97–0.99)			

CI, confidence interval; CCNA, cell cytotoxicity neutralization assay; EIA, enzyme immunoassay; GDH, glutamate dehydrogenase; NAAT, nucleic acid amplification test; TC, toxigenic culture.

Also, none of the studies satisfied all our quality assessment criteria. Notwithstanding these differences, all included studies met the minimal -quite strict- requirements we set. We therefore think that it is justifiable that we calculated summary estimates of sensitivity and specificity, especially because we intended to provide a general overview of test performances of different categories of laboratory assays instead of pointing out one 'best' assay. It is, however, important to realize that test performances of individual assays may have been influenced by the design of included studies analysing these tests. Besides, test characteristics presented here should not be considered unchanging over time and should not be considered fixed characteristics. This is because procedures of commercial assays are sometimes revised to enhance test performance, and also because assays may perform differently among different populations (e.g. high- vs. low-risk patients). Also, in all categories, new assays were marketed. The introduction of newer toxin A/B EIAs leading to a better sensitivity of this category of assays is a good example of the latter.

Table 9. PPV and NPV for different categories of index tests at hypothetical CDI prevalences of 5, 10, 20 and 50%

Test type	CDI prevalence 5%		CDI prevalence 10%		CDI prevalence 20%		CDI prevalence 50%	
	PPV	NPV	PPV	NPV	PPV	NPV	PPV	NPV
Well-type EIA GDH	38	100	54	99	72	98	91	94
Membrane-type EIA GDH	34	100	52	100	71	99	91	98
Well-type EIA toxins A/B	69	99	83	98	91	96	98	87
Membrane-type EIA toxins A/B	81	99	90	98	95	95	99	83
NAAT	46	100	64	100	80	99	94	96

Pooled estimates of sensitivity and specificity compared to cell cytotoxicity neutralization assay were used to calculate the predictive values.

CDI, *Clostridium difficile* infection; EIA, enzyme immunoassay; GDH, glutamate dehydrogenase; NAAT, nucleic acid amplification test; NPV, negative predictive value; PPV, positive predictive value.

On the basis of the review results, PPVs and NPVs were calculated at different hypothetical prevalences of CDI in the tested population. The prevalence of CDI can be seen as the pretest probability of having CDI and would typically be around 5-10% in an endemic setting. (73) At a CDI prevalence of 5%, even the most specific tests (toxin A/B EIAs) would have PPVs of only 69-81%. On the contrary, NPVs would be very high for all index tests. If the prevalence of CDI would rise to 50% among the tested patients, the PPV would consequently raise to 98.8% for the most specific test, but the NPV would drop to 82.5% for the least sensitive tests. Both suboptimal PPV and NPV have implications. A low PPV will result in many patients with false-positive results. These noninfected patients may receive

unnecessary treatment for CDI, and unnecessary isolation precautions may be taken. A low NPV will result in many undetected cases, which may not only have implications for individual patients but also for further transmission of *C. difficile*. It is therefore important to be aware not only of the sensitivity and specificity of an assay but also of the CDI prevalence in the tested population, as the predictive values and hence the clinical utility of the assays depend on them.

The easiest way to diagnose CDI would be to use a single rapid laboratory test that is able to reliably predict disease status. A rapid CDI diagnosis is associated with more prompt CDI treatment and less unnecessarily treated patients. (74) However, two problems arise if the rapid assays are used as stand-alone test for diagnosing CDI. First, as described above, the PPVs of even the most specific tests are inadequate at low disease prevalence. If toxin EIAs were to be used in an endemic situation (CDI prevalence of 5% in the tested population, PPV 81%), an unacceptably high percentage (19%) of patients with a positive test result would not actually have CDI. Second, as the targets identified by the index tests are (just like the targets of the reference test) different from each other, a positive index test does not necessarily indicate a real CDI case. Two of the three categories of index test are not able to differentiate carriers from CDI patients: both GDH EIAs and NAATs do not detect free toxins. Using NAAT as a stand-alone test and relying on clinical symptoms to discern patients with CDI from asymptomatic carriers is not an optimal approach: patients colonized by a toxigenic *C. difficile* strain may very well develop diarrhoea due to other causes, and no specific clinical symptoms exist to differentiate CDI from other causes of diarrhoea. From the above, we conclude that neither GDH EIA nor toxin A/B EIA or NAAT can reliably be used as a stand-alone test to diagnose CDI.

Because no single test is suitable to be used as a stand-alone test, it is best to combine two tests in an algorithm in order to optimize the diagnosis of CDI. The advantage of an algorithm is that tests can be combined in such a way that the percentage of false-positive results can be decreased. This can be done by testing all samples with a first test, then performing reflex testing on samples with a positive first test result only. The first test should be a test that reliably classifies samples with a negative test result as non-CDI; these samples will not be tested further. This first test should therefore be a test with a high NPV (i.e. a highly sensitive test). Thus, in our case, this first test can either be a GDH EIA or NAAT. The choice between these two categories of assays can be made by each individual laboratory. The second test should be a test with a high PPV (i.e. a highly specific test), so that all samples with a positive second test result can reliably be classified as CDI. Toxin

A/B EIAs can very well be used for this purpose, because besides being the most specific tests, these tests also have the advantage of detecting free toxin. Thus, after application of a first sensitive test (GDH EIA or NAAT), the toxin A/B EIA can then be performed as a second step on all samples that tested positive by NAAT or GDH EIA (Fig. 3(a)). Samples with a positive second test result can be classified as CDI likely to be present. However, samples with a first positive test result but a negative toxin A/B EIA need to be clinically evaluated. Among these samples, CDI (with toxin levels below the threshold of detection or a false-negative toxin A/B EIA result) or *C. difficile* carriage is possible. A recent large study tried to establish the optimum diagnostic algorithm for CDI. (47) In this study, 12 420 faecal samples were tested by diverse commercial assays, TC and CCNA. The overall performance of combined tests was superior to individual tests. The combination of a NAAT (Xpert) and toxin A/B EIA (Techlab Tox A/B II) was the optimal algorithm compared to the CCNA test, but the GDH EIA (C. diff Chek-60)-toxin A/B EIA algorithm performed almost identically. (47) These findings can be seen as a validation of our more theoretical approach to establish the best testing strategy, and they endorse the conclusion that NAAT-toxin A/B EIA, or alternatively GDH EIA-toxin A/B EIA, are two of the best algorithms to diagnose CDI (Fig. 3(a)).

An alternative algorithm is to test simultaneously with both a GDH and toxin A/B EIA. An assay is available that includes both these targets in one system (*C. diff* Quik Chek Complete; Techlab), but the sensitivity of the toxin component is unclear and may not be as high as some individual toxin EIAs (Tables 5-7). Samples that test negative for both GDH and toxin A/B can reliably be classified as non-CDI, while samples that test positive for both GDH and toxin A/B can be classified as CDI likely to be present. Samples with a GDH-positive result but that are negative for toxin could undergo reflex testing by NAAT to determine if a toxigenic *C. difficile* strain is present (Fig. 3(b)). Samples with a negative GDH result but that are positive for toxin need to be retested, as this is an invalid result. Only one study evaluating this kind of algorithm and comparing it to a reference test was identified in the literature. (45) In this specific study, samples were screened by *C. diff* Quik Chek Complete, and inconclusive results underwent reflex testing by Illumigene. The overall sensitivity for this algorithm compared to CCNA was 81%, while specificity was reported to be 100%. The overall sensitivity and specificity of this and the aforementioned algorithm depend, however, on the individual assays that are included.

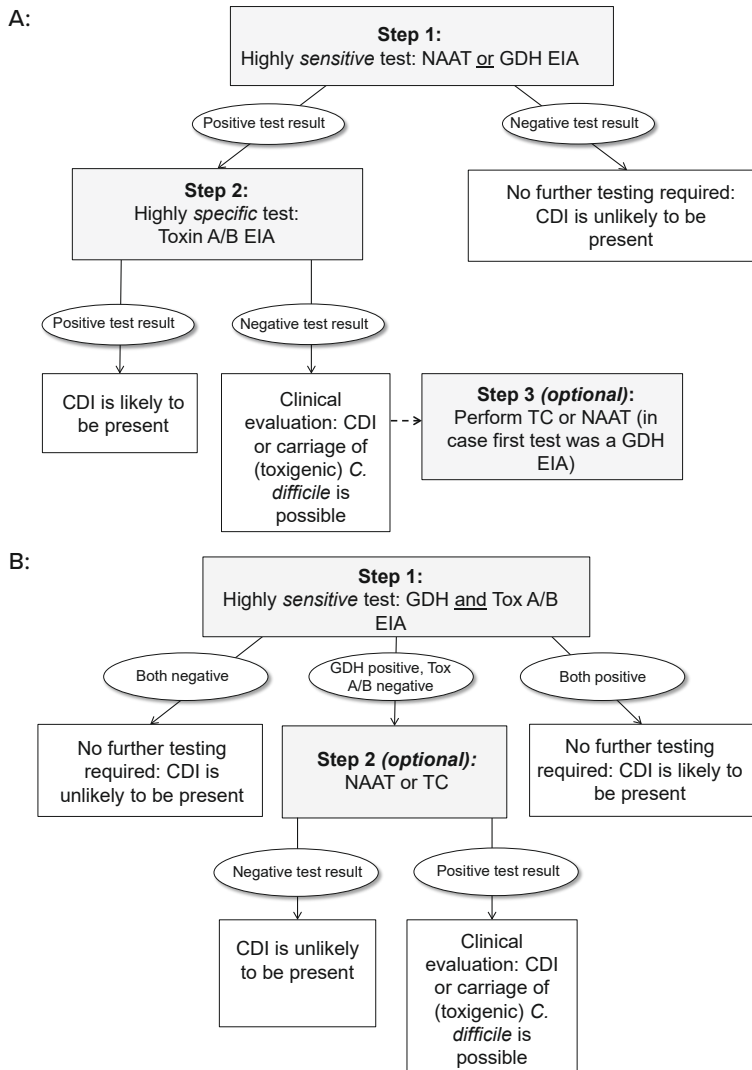


Figure 3. Recommended algorithms for CDI testing. (a) GDH or NAAT–Tox A/B algorithm. (b) GDH and Tox A/B–NAAT/TC algorithm.

CDI, *Clostridium difficile* infection; GDH, glutamate dehydrogenase; NAAT, nucleic acid amplification test; TC, toxigenic culture; Tox A/B, toxin A/B; EIA, enzyme immunoassay.

Although we recommend the use of an algorithm for CDI testing based on two rapid assays, every laboratory should also be able to isolate *C. difficile*, ideally via TC from selected samples, for two reasons. First, TC offers the ability to perform molecular typing and susceptibility testing on recovered isolates from positive samples and can be used for outbreak investigations. (75) Second, samples with a positive GDH EIA and/or NAAT but a

negative toxin A/B EIA may either be samples that tested falsely positive on GDH EIA/NAAT or samples containing *C. difficile*, but without detectable free toxin. To be able to discern between these two conditions, a third-stage reflex test to either a TC or NAAT or GDH (if not yet performed) can be performed on samples with discordant results. For patients with evidence of *C. difficile* but negative toxin A/B EIA, clinical evaluation is needed, and clinical considerations come into play to determine a case as either positive or negative; these patients can either be CDI patients with undetectable toxin levels, or false-negative toxin A/B EIA results or potential carriers of toxigenic *C. difficile*. Although *C. difficile* carriers may play an important role in the spread of the disease (76, 77), the indication for treating these patients for CDI remains controversial. In addition, the need for isolation precautions for these patients remains to be clarified. Therefore, performing TCs on these samples can be of importance for epidemiologic purposes, but it is not yet a prerequisite for patient management.

The decision to treat CDI is ultimately a clinical decision, guided by laboratory results. No tests are infallible, so it may be clinically justified to treat a patient for CDI despite negative test results; treatment should not be withheld on the basis of laboratory tests alone. However, patients with toxin-negative specimens should have alternative diagnoses considered and excluded; provided an adequate testing strategy is followed, most patients with negative results for CDI will truly not have this infection, and thus treatment will be unnecessary.

Besides the question which assay or algorithm should be used for CDI detection, another issue is the number of specimens per patient that should be submitted for testing. Before the introduction of algorithms to diagnose CDI, lack of confidence in the tests for CDI detection (mainly toxin EIAs) led to the practice of multiple sample submission. However, the diagnostic gain of repeat testing within a 7-day period with both toxin A/B EIA and PCR was demonstrated to be very low. (78) If one of the above proposed algorithms is used, then the adequate NPV at low disease prevalence is based on original studies which did not test samples repeatedly by index test and only once by reference test. This adequate NPV indicates that routine submission of multiple samples after a first negative test round has to be discouraged; these samples can reliably be classified as non-CDI.

However, in cases of ongoing clinical suspicion during an endemic situation, the submission of a repeat sample may be justified, as these specific algorithms will have adequate PPVs even in a low-prevalence situation.

In outbreak situations with a higher CDI prevalence in the tested population, the NPV of the algorithm will fall. In such an outbreak situation, submitting a repeat sample in case of ongoing clinical suspicion will be of value, as has been shown for toxin A/B EIA . (79) Testing for cure is not recommended, as patients can shed spores and even toxins of *C. difficile* for a prolonged time after resolution of diarrhoea . (80, 81) The infection can be considered resolved when symptoms of diarrhoea have resolved.

Selection of which of submitted stool samples should be tested for CDI is also important. Recognition of potential CDI cases may be burdensome, as it is increasingly being recognized that CDI is not only acquired in healthcare facilities by patients with well-known risk factors for the disease. In the Netherlands, *C. difficile* was relatively frequent among patients with diarrhoeal complaints in general practice. (82) Community-onset CDI can affect all age groups, and many patients do not have known risk factors. (83, 84) A recent study showed that on a single day in Spain, two of every three CDI episodes were underdiagnosed or misdiagnosed owing to nonsensitive tests (19%) but more importantly to lack of suspicion and request (47.6%). (85) Especially for nonhospitalized patients and younger patients, CDI tests were not requested. (85) This trend was also seen in a study involving almost 500 hospitals in 20 countries across Europe: on two sampling days, 23% of samples with a positive CDI test result were initially missed due to lack of suspicion. (73) Hence, restricting testing to samples with a physician's request for CDI testing will lead to underdiagnosis.

Empirical testing of all unformed stool samples submitted to the laboratory was shown to increase the diagnostic yield. (73, 86) We recommend testing all unformed faecal samples submitted to the laboratory (except samples from children under age 3). In infants, high rates of asymptomatic colonization with both toxigenic and nontoxigenic strains have been described. (87) Even in the case of toxin production, infants rarely develop clinical disease. However, CDI can occur in infants and young children. (88) A recently released policy statement from the American Academy of Pediatrics recommends to test for CDI only if age-specific clinical criteria are met. (14) According to their statement, searching for alternative aetiologies should be performed even in the case of a positive CDI test for children under 3 years of age. Concerning the problematic interpretation of positive test results in this population, we indeed recommend to limit testing of samples from children under age 3 to samples with a physician's request only. Unformed stool samples of children 3 years and older can be managed in the same way as described above.

Clinical signs and symptoms are essential to CDI diagnosis. Therefore, formed stool samples should not be tested for CDI, as these do not meet the clinical criteria of CDI. However, sometimes only solid parts of diarrhoeal faeces may be collected and submitted for *C. difficile* testing. Local protocols therefore need to enable *C. difficile* testing on specific samples to take place. Also, an exception has to be made for patients suspected of CDI who have ileus. In these patients, a rectal swab can be used with adequate sensitivity and specificity for (toxigenic) culture, NAAT or GDH EIA. (89, 90) The use of perirectal swabs for NAAT or GDH EIA testing might also be an alternative in selected patient populations but may depend on the presence of faecal staining of the swab. (89-91) However, the use of (peri)rectal swabs has not been evaluated for toxin EIA, and therefore clinical judgement remains essential in these cases to discern colonized patients from patients with CDI.

Recommendations

Sample selection

- We recommend that CDI testing should not be limited to samples with a specific physician's request. (Strong recommendation, high-quality evidence)
- We suggest that at least all submitted unformed stool samples from patients 3 years or older should be tested for CDI. (Weak recommendation, low-quality evidence)
- We suggest to limit testing of samples from children under age 3 to samples with a physician's request only. (Weak recommendation, low-quality evidence)
- Formed stool samples should not be tested for CDI (except in case of paralytic ileus). (Good practice statement)
- In patients suspected of ileus, a rectal swab can be used for (toxigenic) culture, NAAT or GDH EIA. (Strong recommendation, moderate-quality evidence)

Testing protocol

- The diagnosis of CDI should be based on clinical signs and symptoms in combination with laboratory tests. Decision for treatment for CDI is a clinical decision and may be justified even if all laboratory tests are negative. (Good practice statement)
- We recommend against the use of a single rapid test as a stand-alone test due to inadequate PPV in an endemic situation. (Strong recommendation, moderate-quality evidence)

- We recommend the use of a 2-step algorithm (Fig. 3(A)). (Strong recommendation, moderate-quality evidence)
- This algorithm should start with either NAAT or GDH EIA. Samples with a negative first test result can be reported as negative. (Strong recommendation, moderate-quality evidence)
- Samples with a positive first test result should be tested further with a toxin A/B EIA. Samples with a positive second test results can be reported as CDI-positive. (Strong recommendation, moderate-quality evidence)
- An alternative algorithm is to screen samples with both a GDH and toxin A/B EIA (Fig. 3(B)). Samples with concordant positive or negative results can be reported as such. Samples with a negative GDH result but positive for toxin need to be retested as this is an invalid result. (Strong recommendation, moderate-quality evidence)
- Samples with a positive first test result and negative second test result (Fig. 3(A)) and samples with a GDH-positive test result but negative toxin A/B test result (Fig. 3(B)) may represent samples with CDI or *C. difficile* carriage and may optionally be tested with TC or NAAT (if not performed yet). (Weak recommendation, moderate-quality evidence)
- We recommend to perform TC and molecular typing of recovered isolates in case of outbreak situations. (Good practice statement)

Repeated testing

- Repeated testing after a first positive sample during the same diarrhoeal episode is not recommended in an endemic situation. (Strong recommendation, moderate-quality evidence)
- Repeated testing after a first negative sample during the same diarrhoeal episode may be useful in selected cases with ongoing clinical suspicion during an epidemic situation or in cases with high clinical suspicion during endemic situations. (Strong recommendation, moderate-quality evidence)
- A test of cure is not recommended. (Good practice statement)

Acknowledgements

Members of the Executive Committee are as follows: M. J. T. Crobach, Department of Medical Microbiology, Centre for Infectious Diseases, Leiden University Medical Center, Leiden, The Netherlands; T. Planche, Department of Medical Microbiology, St George's Hospital, London, UK; C. Eckert and F. Barbut, ESGCD members, National Reference Laboratory for

Chapter 3

Clostridium difficile, Paris, France; O. M. Dekkers, Departments of Clinical Epidemiology and Internal Medicine, Leiden University Medical Center, Leiden, The Netherlands, and Department of Clinical Epidemiology, Aarhus University, Aarhus, Denmark; E. M. Terveer, Department of Medical Microbiology, Centre for Infectious Diseases, Leiden University Medical Center, Leiden, The Netherlands; M. H. Wilcox, ESGCD member, Department of Microbiology, Leeds Teaching Hospitals & University of Leeds, Leeds, UK; E. J. Kuijper (chair), ESGCD member, Department of Medical Microbiology, Centre for Infectious Diseases, Leiden University Medical Center, Leiden, The Netherlands.

Members of the Drafting Group are as follows: F. Allerberger, ESGCD member, Austrian Agency for Health and Food Safety (AGES), Vienna, Austria; L. von Müller, ESGCD member, Institut für Labormedizin, Mikrobiologie und Hygiene (LMH), Christophorus-Kliniken GmbH, Coesfeld, and Advisory Laboratory for *Clostridium difficile*, Homburg/Saar, Germany; F. Fitzpatrick, ESGCD member, Department of Clinical Microbiology, Microbiology Laboratory, Beaumont Hospital, Ireland; R. Frei, ESGCD member, Division of Clinical Microbiology, University Hospital Basel, Basel, Switzerland; S. Mentula, ESGCD member, Bacterial Infections Unit, National Institute for Health and Welfare, Helsinki, Finland; J. E. Coia, ESGCD member, Department of Clinical Microbiology, Glasgow Royal Infirmary, Glasgow, UK; T. Nore'n, ESGCD member, Department of Infection Control, Örebro University Hospital, Örebro, Sweden; J. van Broeck and M. Delmee, ESGCD members, Université Catholique de Louvain, Microbiology Unit, Brussels, Belgium.

Transparency Declaration

All authors report no conflicts of interest relevant to this article.

References

1. Crobach MJ, Dekkers OM, Wilcox MH, Kuijper EJ. European Society of Clinical Microbiology and Infectious Diseases (ESCMID): data review and recommendations for diagnosing *Clostridium difficile*-infection (CDI). *Clinical microbiology and infection : the official publication of the European Society of Clinical Microbiology and Infectious Diseases*. 2009;15(12):1053-66.
2. Planche T, Wilcox M. Reference assays for *Clostridium difficile* infection: one or two gold standards? *J Clin Pathol*. 2011;64(1):1-5.
3. Delmee M. Laboratory diagnosis of *Clostridium difficile* disease. *Clinical microbiology and infection : the official publication of the European Society of Clinical Microbiology and Infectious Diseases*. 2001;7(8):411-6.
4. Hink T, Burnham CA, Dubberke ER. A systematic evaluation of methods to optimize culture-based recovery of *Clostridium difficile* from stool specimens. *Anaerobe*. 2013;19:39-43.
5. Borriello SP, Honour P. Simplified procedure for the routine isolation of *Clostridium difficile* from faeces. *J Clin Pathol*. 1981;34(10):1124-7.
6. Burnham CA, Carroll KC. Diagnosis of *Clostridium difficile* infection: an ongoing conundrum for clinicians and for clinical laboratories. *Clinical microbiology reviews*. 2013;26(3):604-30.
7. Lalande V, Barrault L, Wadel S, Eckert C, Petit JC, Barbut F. Evaluation of a loop-mediated isothermal amplification assay for diagnosis of *Clostridium difficile* infections. *J Clin Microbiol*. 2011;49(7):2714-6.
8. Eckert C, Holscher E, Petit A, Lalande V, Barbut F. Molecular test based on isothermal helicase-dependent amplification for detection of the *Clostridium difficile* toxin A gene. *J Clin Microbiol*. 2014;52(7):2386-9.
9. Nyč O, Pituch H, Matějková J, Obuch-Woszczatynski P, Kuijper EJ. *Clostridium difficile* PCR ribotype 176 in the Czech Republic and Poland. *Lancet*. 2011;377(9775):1407.
10. Whiting PF, Rutjes AW, Westwood ME, Mallett S, Deeks JJ, Reitsma JB, et al. QUADAS-2: a revised tool for the quality assessment of diagnostic accuracy studies. *Ann Intern Med*. 2011;155(8):529-36.
11. Cohen SH, Gerding DN, Johnson S, Kelly CP, Loo VG, McDonald LC, et al. Clinical practice guidelines for *Clostridium difficile* infection in adults: 2010 update by the society for healthcare epidemiology of America (SHEA) and the infectious diseases society of America (IDSA). *Infect Control Hosp Epidemiol*. 2010;31(5):431-55.
12. Cheng AC, Ferguson JK, Richards MJ, Robson JM, Gilbert GL, McGregor A, et al. Australasian Society for Infectious Diseases guidelines for the diagnosis and treatment of *Clostridium difficile* infection. *Med J Aust*. 2011;194(7):353-8.
13. Surawicz CM, Brandt LJ, Binion DG, Ananthakrishnan AN, Curry SR, Gilligan PH, et al. Guidelines for diagnosis, treatment, and prevention of *Clostridium difficile* infections. *Am J Gastroenterol*. 2013;108(4):478-98; quiz 99.
14. Schutze GE, Willoughby RE. *Clostridium difficile* infection in infants and children. *Pediatrics*. 2013;131(1):196-200.

15. Updated guidance on the diagnosis and reporting of *Clostridium difficile*. . In: NHS, editor. London.
16. Brouwers MC, Kho ME, Browman GP, Burgers JS, Cluzeau F, Feder G, et al. AGREE II: advancing guideline development, reporting and evaluation in health care. *J Clin Epidemiol*. 2010;63(12):1308-11.
17. Schünemann HJ, Oxman AD, Brozek J, Glasziou P, Jaeschke R, Vist GE, et al. Grading quality of evidence and strength of recommendations for diagnostic tests and strategies. *Bmj*. 2008;336(7653):1106-10.
18. Guyatt GH, Schünemann HJ, Djulbegovic B, Akl EA. Guideline panels should not GRADE good practice statements. *J Clin Epidemiol*. 2015;68(5):597-600.
19. Barkin JA, Nandi N, Miller N, Grace A, Barkin JS, Sussman DA. Superiority of the DNA amplification assay for the diagnosis of *C. difficile* infection: a clinical comparison of fecal tests. *Dig Dis Sci*. 2012;57(10):2592-9.
20. Berry N, Sewell B, Jafri S, Puli C, Vagia S, Lewis AM, et al. Real-time polymerase chain reaction correlates well with clinical diagnosis of *Clostridium difficile* infection. *J Hosp Infect*. 2014;87(2):109-14.
21. Bruins MJ, Verbeek E, Wallinga JA, Bruijnesteijn van Coppenraet LE, Kuijper EJ, Bloembergen P. Evaluation of three enzyme immunoassays and a loop-mediated isothermal amplification test for the laboratory diagnosis of *Clostridium difficile* infection. *Eur J Clin Microbiol Infect Dis*. 2012;31(11):3035-9.
22. Buchan BW, Mackey TL, Daly JA, Alger G, Denys GA, Peterson LR, et al. Multicenter clinical evaluation of the portrait toxigenic *C. difficile* assay for detection of toxigenic *Clostridium difficile* strains in clinical stool specimens. *J Clin Microbiol*. 2012;50(12):3932-6.
23. Calderaro A, Buttrini M, Martinelli M, Gorrini C, Montecchini S, Medici MC, et al. Comparative analysis of different methods to detect *Clostridium difficile* infection. *New Microbiol*. 2013;36(1):57-63.
24. Carroll KC, Buchan BW, Tan S, Stamper PD, Riebe KM, Pancholi P, et al. Multicenter evaluation of the Verigene *Clostridium difficile* nucleic acid assay. *J Clin Microbiol*. 2013;51(12):4120-5.
25. de Boer RF, Wijma JJ, Schuurman T, Moedt J, Dijk-Alberts BG, Ott A, et al. Evaluation of a rapid molecular screening approach for the detection of toxigenic *Clostridium difficile* in general and subsequent identification of the *tcdC* Δ 117 mutation in human stools. *J Microbiol Methods*. 2010;83(1):59-65.
26. de Jong E, de Jong AS, Bartels CJ, van der Rijt-van den Biggelaar C, Melchers WJ, Sturm PD. Clinical and laboratory evaluation of a real-time PCR for *Clostridium difficile* toxin A and B genes. *Eur J Clin Microbiol Infect Dis*. 2012;31(9):2219-25.
27. Eastwood K, Else P, Charlett A, Wilcox M. Comparison of nine commercially available *Clostridium difficile* toxin detection assays, a real-time PCR assay for *C. difficile* *tcdB*, and a glutamate dehydrogenase detection assay to cytotoxin testing and cytotoxigenic culture methods. *J Clin Microbiol*. 2009;47(10):3211-7.
28. Fenner L, Widmer AF, Goy G, Rudin S, Frei R. Rapid and reliable diagnostic algorithm for detection of *Clostridium difficile*. *J Clin Microbiol*. 2008;46(1):328-30.

29. Hart J, Putsathit P, Knight DR, Sammels L, Riley TV, Keil A. Clostridium difficile infection diagnosis in a paediatric population: comparison of methodologies. *Eur J Clin Microbiol Infect Dis.* 2014;33(9):1555-64.
30. Hirvonen JJ, Mentula S, Kaukoranta SS. Evaluation of a new automated homogeneous PCR assay, GenomEra C. difficile, for rapid detection of Toxigenic Clostridium difficile in fecal specimens. *J Clin Microbiol.* 2013;51(9):2908-12.
31. Huang H, Weintraub A, Fang H, Nord CE. Comparison of a commercial multiplex real-time PCR to the cell cytotoxicity neutralization assay for diagnosis of clostridium difficile infections. *J Clin Microbiol.* 2009;47(11):3729-31.
32. Jacobs J, Rudensky B, Dresner J, Berman A, Sonnenblick M, van Dijk Y, et al. Comparison of four laboratory tests for diagnosis of Clostridium difficile-associated diarrhea. *Eur J Clin Microbiol Infect Dis.* 1996;15(7):561-6.
33. Kawada M, Annaka M, Kato H, Shibasaki S, Hikosaka K, Mizuno H, et al. Evaluation of a simultaneous detection kit for the glutamate dehydrogenase antigen and toxin A/B in feces for diagnosis of Clostridium difficile infection. *J Infect Chemother.* 2011;17(6):807-11.
34. Kim H, Jeong SH, Kim M, Lee Y, Lee K. Detection of Clostridium difficile toxin A/B genes by multiplex real-time PCR for the diagnosis of C. difficile infection. *J Med Microbiol.* 2012;61(Pt 2):274-7.
35. Kim H, Kim WH, Kim M, Jeong SH, Lee K. Evaluation of a rapid membrane enzyme immunoassay for the simultaneous detection of glutamate dehydrogenase and toxin for the diagnosis of Clostridium difficile infection. *Ann Lab Med.* 2014;34(3):235-9.
36. Larson AM, Fung AM, Fang FC. Evaluation of tcdB real-time PCR in a three-step diagnostic algorithm for detection of toxigenic Clostridium difficile. *J Clin Microbiol.* 2010;48(1):124-30.
37. Le Guern R, Herwegh S, Grandbastien B, Courcol R, Wallet F. Evaluation of a new molecular test, the BD Max Cdiff, for detection of toxigenic Clostridium difficile in fecal samples. *J Clin Microbiol.* 2012;50(9):3089-90.
38. Leitner E, Einetter M, Grisold AJ, Marth E, Feierl G. Evaluation of the BD MAX Cdiff assay for the detection of the toxin B gene of Clostridium difficile out of faecal specimens. *Diagn Microbiol Infect Dis.* 2013;76(3):390-1.
39. Massey V, Gregson DB, Chagla AH, Storey M, John MA, Hussain Z. Clinical usefulness of components of the Triage immunoassay, enzyme immunoassay for toxins A and B, and cytotoxin B tissue culture assay for the diagnosis of Clostridium difficile diarrhea. *Am J Clin Pathol.* 2003;119(1):45-9.
40. Mattner F, Winterfeld I, Mattner L. Diagnosing toxigenic Clostridium difficile: new confidence bounds show culturing increases sensitivity of the toxin A/B enzyme immunoassay and refute gold standards. *Scand J Infect Dis.* 2012;44(8):578-85.
41. Musher DM, Manhas A, Jain P, Nuila F, Waqar A, Logan N, et al. Detection of Clostridium difficile toxin: comparison of enzyme immunoassay results with results obtained by cytotoxicity assay. *J Clin Microbiol.* 2007;45(8):2737-9.
42. Norén T, Alriksson I, Andersson J, Akerlund T, Unemo M. Rapid and sensitive loop-mediated isothermal amplification test for Clostridium difficile detection challenges cytotoxin B cell test and culture as gold standard. *J Clin Microbiol.* 2011;49(2):710-1.

43. Novak-Weekley SM, Marlowe EM, Miller JM, Cumpio J, Nomura JH, Vance PH, et al. Clostridium difficile testing in the clinical laboratory by use of multiple testing algorithms. *J Clin Microbiol.* 2010;48(3):889-93.
44. O'Connor D, Hynes P, Cormican M, Collins E, Corbett-Feeney G, Cassidy M. Evaluation of methods for detection of toxins in specimens of feces submitted for diagnosis of Clostridium difficile-associated diarrhea. *J Clin Microbiol.* 2001;39(8):2846-9.
45. Ota KV, McGowan KL. Clostridium difficile testing algorithms using glutamate dehydrogenase antigen and C. difficile toxin enzyme immunoassays with C. difficile nucleic acid amplification testing increase diagnostic yield in a tertiary pediatric population. *J Clin Microbiol.* 2012;50(4):1185-8.
46. Pancholi P, Kelly C, Raczkowski M, Balada-Llasat JM. Detection of toxigenic Clostridium difficile: comparison of the cell culture neutralization, Xpert C. difficile, Xpert C. difficile/Epi, and Illumigene C. difficile assays. *J Clin Microbiol.* 2012;50(4):1331-5.
47. Planche TD, Davies KA, Coen PG, Finney JM, Monahan IM, Morris KA, et al. Differences in outcome according to Clostridium difficile testing method: a prospective multicentre diagnostic validation study of C difficile infection. *The Lancet Infectious Diseases.* 2013;13(11):936-45.
48. Qutub MO, AlBaz N, Hawken P, Anoos A. Comparison between the two-step and the three-step algorithms for the detection of toxigenic Clostridium difficile. *Indian J Med Microbiol.* 2011;29(3):293-6.
49. Reller ME, Lema CA, Perl TM, Cai M, Ross TL, Speck KA, et al. Yield of stool culture with isolate toxin testing versus a two-step algorithm including stool toxin testing for detection of toxigenic Clostridium difficile. *J Clin Microbiol.* 2007;45(11):3601-5.
50. Reller ME, Alcabasa RC, Lema CA, Carroll KC. Comparison of two rapid assays for Clostridium difficile Common antigen and a C difficile toxin A/B assay with the cell culture neutralization assay. *Am J Clin Pathol.* 2010;133(1):107-9.
51. Shin BM, Lee EJ, Kuak EY, Yoo SJ. Comparison of VIDAS CDAB and CDA immunoassay for the detection of Clostridium difficile in a tcdA- tcdB+ C. difficile prevalent area. *Anaerobe.* 2009;15(6):266-9.
52. Shin BM, Kuak EY, Lee EJ, Songer JG. Algorithm combining toxin immunoassay and stool culture for diagnosis of Clostridium difficile infection. *J Clin Microbiol.* 2009;47(9):2952-6.
53. Shin BM, Mun SJ, Yoo SJ, Kuak EY. Comparison of BD GeneOhm Cdiff and Seegene Seeplex ACE PCR assays using toxigenic Clostridium difficile culture for direct detection of tcdB from stool specimens. *J Clin Microbiol.* 2012;50(11):3765-7.
54. Shin S, Kim M, Kim M, Lim H, Kim H, Lee K, et al. Evaluation of the Xpert Clostridium difficile assay for the diagnosis of Clostridium difficile infection. *Ann Lab Med.* 2012;32(5):355-8.
55. Sloan LM, Duresko BJ, Gustafson DR, Rosenblatt JE. Comparison of real-time PCR for detection of the tcdC gene with four toxin immunoassays and culture in diagnosis of Clostridium difficile infection. *J Clin Microbiol.* 2008;46(6):1996-2001.

56. Snell H, Ramos M, Longo S, John M, Hussain Z. Performance of the TechLab C. DIFF CHEK-60 enzyme immunoassay (EIA) in combination with the C. difficile Tox A/B II EIA kit, the Triage C. difficile panel immunoassay, and a cytotoxin assay for diagnosis of *Clostridium difficile*-associated diarrhea. *J Clin Microbiol.* 2004;42(10):4863-5.
57. Soh YS, Yang JJ, You E, La Jeon Y, Kim MJ, Nam YS, et al. Comparison of two molecular methods for detecting toxigenic *Clostridium difficile*. *Ann Clin Lab Sci.* 2014;44(1):27-31.
58. Stamper PD, Babiker W, Alcabasa R, Aird D, Wehrin J, Ikpeama I, et al. Evaluation of a new commercial TaqMan PCR assay for direct detection of the *clostridium difficile* toxin B gene in clinical stool specimens. *J Clin Microbiol.* 2009;47(12):3846-50.
59. Stamper PD, Alcabasa R, Aird D, Babiker W, Wehrin J, Ikpeama I, et al. Comparison of a commercial real-time PCR assay for *tcdB* detection to a cell culture cytotoxicity assay and toxigenic culture for direct detection of toxin-producing *Clostridium difficile* in clinical samples. *J Clin Microbiol.* 2009;47(2):373-8.
60. Staneck JL, Weckbach LS, Allen SD, Siders JA, Gilligan PH, Coppitt G, et al. Multicenter evaluation of four methods for *Clostridium difficile* detection: ImmunoCard C. *difficile*, cytotoxin assay, culture, and latex agglutination. *J Clin Microbiol.* 1996;34(11):2718-21.
61. Swindells J, Brenwald N, Reading N, Oppenheim B. Evaluation of diagnostic tests for *Clostridium difficile* infection. *J Clin Microbiol.* 2010;48(2):606-8.
62. Tenover FC, Novak-Weekley S, Woods CW, Peterson LR, Davis T, Schreckenberger P, et al. Impact of strain type on detection of toxigenic *Clostridium difficile*: comparison of molecular diagnostic and enzyme immunoassay approaches. *J Clin Microbiol.* 2010;48(10):3719-24.
63. Terhes G, Urbán E, Sóki J, Nacsá E, Nagy E. Comparison of a rapid molecular method, the BD GeneOhm Cdiff assay, to the most frequently used laboratory tests for detection of toxin-producing *Clostridium difficile* in diarrheal feces. *J Clin Microbiol.* 2009;47(11):3478-81.
64. Ticehurst JR, Aird DZ, Dam LM, Borek AP, Hargrove JT, Carroll KC. Effective detection of toxigenic *Clostridium difficile* by a two-step algorithm including tests for antigen and cytotoxin. *J Clin Microbiol.* 2006;44(3):1145-9.
65. Turgeon DK, Novicki TJ, Quick J, Carlson L, Miller P, Ulness B, et al. Six rapid tests for direct detection of *Clostridium difficile* and its toxins in fecal samples compared with the fibroblast cytotoxicity assay. *J Clin Microbiol.* 2003;41(2):667-70.
66. van den Berg RJ, Bruijnesteijn van Coppenraet LS, Gerritsen HJ, Endtz HP, van der Vorm ER, Kuijper EJ. Prospective multicenter evaluation of a new immunoassay and real-time PCR for rapid diagnosis of *Clostridium difficile*-associated diarrhea in hospitalized patients. *J Clin Microbiol.* 2005;43(10):5338-40.
67. van den Berg RJ, Vaessen N, Endtz HP, Schölin T, van der Vorm ER, Kuijper EJ. Evaluation of real-time PCR and conventional diagnostic methods for the detection of *Clostridium difficile*-associated diarrhoea in a prospective multicentre study. *J Med Microbiol.* 2007;56(Pt 1):36-42.
68. Vanpoucke H, De Baere T, Claeys G, Vaneechoutte M, Verschraegen G. Evaluation of six commercial assays for the rapid detection of *Clostridium difficile* toxin and/or antigen in stool specimens. *Clinical microbiology and infection : the official publication of the European Society of Clinical Microbiology and Infectious Diseases.* 2001;7(2):55-64.

69. Viala C, Le Monnier A, Maataoui N, Rousseau C, Collignon A, Poilane I. Comparison of commercial molecular assays for toxigenic *Clostridium difficile* detection in stools: BD GeneOhm Cdiff, XPert C. *difficile* and illumigene C. *difficile*. *J Microbiol Methods*. 2012;90(2):83-5.
70. Walkty A, Lagacé-Wiens PR, Manickam K, Adam H, Pieroni P, Hoban D, et al. Evaluation of an algorithmic approach in comparison with the Illumigene assay for laboratory diagnosis of *Clostridium difficile* infection. *J Clin Microbiol*. 2013;51(4):1152-7.
71. Wren MW, Kinson R, Sivapalan M, Shemko M, Shetty NR. Detection of *Clostridium difficile* infection: a suggested laboratory diagnostic algorithm. *Br J Biomed Sci*. 2009;66(4):175-9.
72. Zheng L, Keller SF, Lyerly DM, Carman RJ, Genheimer CW, Gleaves CA, et al. Multicenter evaluation of a new screening test that detects *Clostridium difficile* in fecal specimens. *J Clin Microbiol*. 2004;42(8):3837-40.
73. Davies KA, Longshaw CM, Davis GL, Bouza E, Barbut F, Barna Z, et al. Underdiagnosis of *Clostridium difficile* across Europe: the European, multicentre, prospective, biannual, point-prevalence study of *Clostridium difficile* infection in hospitalised patients with diarrhoea (EUCLID). *Lancet Infect Dis*. 2014;14(12):1208-19.
74. Barbut F, Surgers L, Eckert C, Visseaux B, Cuingnet M, Mesquita C, et al. Does a rapid diagnosis of *Clostridium difficile* infection impact on quality of patient management? *Clinical microbiology and infection : the official publication of the European Society of Clinical Microbiology and Infectious Diseases*. 2014;20(2):136-44.
75. Goorhuis A, Debast SB, Dutilh JC, van Kinschot CM, Harmanus C, Cannegieter SC, et al. Type-specific risk factors and outcome in an outbreak with 2 different *Clostridium difficile* types simultaneously in 1 hospital. *Clinical infectious diseases : an official publication of the Infectious Diseases Society of America*. 2011;53(9):860-9.
76. Curry SR, Muto CA, Schlackman JL, Pasculle AW, Shutt KA, Marsh JW, et al. Use of multilocus variable number of tandem repeats analysis genotyping to determine the role of asymptomatic carriers in *Clostridium difficile* transmission. *Clinical infectious diseases : an official publication of the Infectious Diseases Society of America*. 2013;57(8):1094-102.
77. Eyre DW, Griffiths D, Vaughan A, Golubchik T, Acharya M, O'Connor L, et al. Asymptomatic *Clostridium difficile* colonisation and onward transmission. *PLoS One*. 2013;8(11):e78445.
78. Aichinger E, Schleck CD, Harmsen WS, Nyre LM, Patel R. Nonutility of repeat laboratory testing for detection of *Clostridium difficile* by use of PCR or enzyme immunoassay. *J Clin Microbiol*. 2008;46(11):3795-7.
79. Debast SB, van Kregten E, Oskam KM, van den Berg T, Van den Berg RJ, Kuijper EJ. Effect on diagnostic yield of repeated stool testing during outbreaks of *Clostridium difficile*-associated disease. *Clinical microbiology and infection : the official publication of the European Society of Clinical Microbiology and Infectious Diseases*. 2008;14(6):622-4.
80. Sethi AK, Al-Nassir WN, Nerandzic MM, Bobulsky GS, Donskey CJ. Persistence of skin contamination and environmental shedding of *Clostridium difficile* during and after treatment of *C. difficile* infection. *Infect Control Hosp Epidemiol*. 2010;31(1):21-7.

81. Wenisch C, Parschalk B, Hasenhundl M, Hirschl AM, Graninger W. Comparison of vancomycin, teicoplanin, metronidazole, and fusidic acid for the treatment of *Clostridium difficile*-associated diarrhea. *Clinical infectious diseases : an official publication of the Infectious Diseases Society of America*. 1996;22(5):813-8.
82. Hensgens MP, Dekkers OM, Demeulemeester A, Buiting AG, Bloembergen P, van Benthem BH, et al. Diarrhoea in general practice: when should a *Clostridium difficile* infection be considered? Results of a nested case-control study. *Clinical microbiology and infection : the official publication of the European Society of Clinical Microbiology and Infectious Diseases*. 2014;20(12):O1067-74.
83. Bauer MP, Veenendaal D, Verhoef L, Bloembergen P, van Dissel JT, Kuijper EJ. Clinical and microbiological characteristics of community-onset *Clostridium difficile* infection in The Netherlands. *Clinical microbiology and infection : the official publication of the European Society of Clinical Microbiology and Infectious Diseases*. 2009;15(12):1087-92.
84. Wilcox MH, Mooney L, Bendall R, Settle CD, Fawley WN. A case-control study of community-associated *Clostridium difficile* infection. *J Antimicrob Chemother*. 2008;62(2):388-96.
85. Alcalá L, Martín A, Marin M, Sánchez-Somolinos M, Catalan P, Peláez T, et al. The undiagnosed cases of *Clostridium difficile* infection in a whole nation: where is the problem? *Clinical microbiology and infection : the official publication of the European Society of Clinical Microbiology and Infectious Diseases*. 2012;18(7):E204-13.
86. Reigadas E, Alcalá L, Marin M, Burillo A, Muñoz P, Bouza E. Missed diagnosis of *Clostridium difficile* infection; a prospective evaluation of unselected stool samples. *J Infect*. 2015;70(3):264-72.
87. Bryant K, McDonald LC. *Clostridium difficile* infections in children. *Pediatr Infect Dis J*. 2009;28(2):145-6.
88. Wendt JM, Cohen JA, Mu Y, Dumyati GK, Dunn JR, Holzbauer SM, et al. *Clostridium difficile* infection among children across diverse US geographic locations. *Pediatrics*. 2014;133(4):651-8.
89. Rogers DS, Kundrapu S, Sunkesula VC, Donskey CJ. Comparison of perirectal versus rectal swabs for detection of asymptomatic carriers of toxigenic *Clostridium difficile*. *J Clin Microbiol*. 2013;51(10):3421-2.
90. McFarland LV, Coyle MB, Kremer WH, Stamm WE. Rectal swab cultures for *Clostridium difficile* surveillance studies. *J Clin Microbiol*. 1987;25(11):2241-2.
91. Kundrapu S, Sunkesula VC, Jury LA, Sethi AK, Donskey CJ. Utility of perirectal swab specimens for diagnosis of *Clostridium difficile* infection. *Clinical infectious diseases : an official publication of the Infectious Diseases Society of America*. 2012;55(11):1527-30.