

Certificate of Analysis for NR-49324

Clostridium difficile, Isolate 20121190

Catalog No. NR-49324

Product Description: Clostridium difficile (C. difficile; also referred to as Peptoclostridium difficile), isolate 20121190 was obtained from the stool of an elderly female patient with a healthcare-associated (HA) C. difficile infection in midwestern USA, in 2011. Isolate 20121190 was deposited as PCR ribotype 126, North American pulsed-field gel electrophoresis type 12 (NAP12), containing tcdA, tcdB and tcdC (with 18 base pair deletion) of the PaLoc operon. This isolate is reported to be negative for the C. difficile binary toxin (CDT).

Lot¹: 63719917 Manufacturing Date: 21OCT2015

TEST	SPECIFICATIONS	RESULTS
Phenotypic Analysis		
Cellular morphology	Gram-positive rods	Gram-positive rods
Colony morphology ²	Report results	Irregular, flat, undulate, rough and gray (Figure 1)
Hemolysis ²	Non-hemolytic	Non-hemolytic
Motility (wet mount)	Report results	Motile
Biochemical tests:		
Esculin hydrolysis ³	Positive	Positive
Gelatin hydrolysis ³	Positive	Positive
VITEK® MS (MALDI-TOF)	Consistent with C. difficile	Consistent with <i>C. difficile</i> (99.9%)
Genotypic Analysis		
Sequencing of 16S ribosomal RNA gene	≥ 99% sequence identity to	99.2% sequence identity to
(~ 1320 base pairs)	C. difficile type strain	AB075770
PCR Assay of Extracted DNA		
Presence of <i>C. difficile</i> -specific genes ⁴		
Triose phosphate isomerase (tpi)	~ 230 base pair amplicon	~ 230 base pair amplicon
Presence of toxin genes ^{4,5}		
cdtB	No amplicon	No amplicon
tcdA (wild type)	~ 370 base pair amplicon	~ 370 base pair amplicon
tcdA (partial deletion)	No amplicon	No amplicon
tcdB	~ 160 base pair amplicon	~ 160 base pair amplicon
Purity (post-freeze)		
Anaerobic growth ⁶	Growth consistent with expected	Growth consistent with expected
7	colony morphology	colony morphology
Aerobic growth ⁷	No growth	No growth
Viability (post-freeze) ²	Growth	Growth

¹NR-49324 was produced by inoculation of the deposited material into Modified Reinforced Clostridial medium and incubated for 1 day at 37°C in an anaerobic atmosphere (< 0.5% O₂; Remel™ Anaero Pack-Anaero™ R681001). The material from the initial growth was passaged in Modified Reinforced Clostridial medium for 1 day at 37°C in an anaerobic atmosphere to produce this lot.

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²2 days at 37°C in an anaerobic atmosphere on Tryptic Soy agar with 5% defibrinated sheep blood

³Tests were assessed after 7 days at 37°C in an anaerobic atmosphere. The gelatin tube was placed at 4°C for one hour prior to result determination.

⁴Lemee, L., et al. "Multiplex PCR Targeting *tpi* (Triose Phosphate Isomerase), *tcdA* (Toxin A), and *tcdB* (Toxin B) Genes for Toxigenic Culture of *Clostridium difficile*." J. Clin. Microbiol. 42 (2004): 5710-5714. PubMed: 15583303.

⁵Antikainen, J., et al. "Detection of Virulence Genes of Clostridium difficile by Multiplex PCR." <u>APMIS.</u> 117 (2009): 607-613. PubMed: 19664132.

⁶Purity of this lot was assessed for 7 days at 37°C in an anaerobic atmosphere on Tryptic Soy agar with 5% defibrinated sheep blood.

⁷Purity of this lot was assessed for 7 days at 37°C in an aerobic atmosphere with 5% CO₂ on Tryptic Soy agar with 5% defibrinated sheep blood.



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Figure 1: Colony Morphology



Date: 18 FEB 2016

Signature:

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