**sodC-Based Real-Time PCR for Detection of *Neisseria meningitidis***

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**Abstract**

Real-time PCR (rt-PCR) is a widely used molecular method for detection of *Neisseria meningitidis* (Nm). Several rt-PCR assays for Nm target the capsule transport gene, ctrA. However, over 16% of meningococcal carriage isolates lack ctrA, rendering this target gene ineffective at identification of this sub-population of meningococcal isolates. The Cu-Zn superoxide dismutase gene, sodC, is found in Nm but not in other *Neisseria* species. To better identify Nm, regardless of capsule genotype or expression status, a sodC-based TaqMan rt-PCR assay was developed and validated. Standard curves revealed an average lower limit of detection of 73 genomes per reaction at cycle threshold (Ct) value of 35, with 100% average reaction efficiency and an average R2 of 0.9925. 99.7% (624/626) of Nm isolates tested were sodC-positive, with a range of average Ct values from 13.0 to 29.5. The mean sodC Ct value of these Nm isolates was 17.6±2.2 (±SD). Of the 626 Nm tested, 178 were nongroupable (NG) ctrA-negative Nm isolates, and 98.9% (176/178) of these were detected by sodC rt-PCR. The assay was 100% specific, with all 244 non-Nm isolates testing negative. Of 157 clinical specimens tested, sodC detected 25/157 Nm or 4 additional specimens compared to ctrA and 24 more than culture. Among 582 carriage specimens, sodC detected Nm in 1 more than ctrA and in 4 more than culture. This sodC rt-PCR assay is a highly sensitive and specific method for detection of Nm, especially in carriage studies where many meningococcal isolates lack capsule genes.

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**Introduction**

*Neisseria meningitidis* (Nm) is the etiologic agent of epidemic bacterial meningitis and rapidly fatal sepsis throughout the world. Many clinical, reference, and research laboratories must be able to rapidly detect Nm either from patients with invasive disease or from asymptomatic carriers. Bacterial isolates or clinical specimens may be sent to the laboratory from patients with suspected meningitis, while isolates or swab eluates may be the specimens that come to the laboratory from possible carriers.

Common techniques employed for the identification of Nm include biochemical tests, slide agglutination serogrouping (SASG) [1,2], and the polymerase chain reaction (PCR) [3–6]. Chromogenic biochemical tests and SASG can be subjective, sometimes complicating species identification [3,7].

Unlike biochemical tests and SASG, PCR does not require viable bacteria and can be used to identify and characterize even nongroupable (NG) meningococci. Additionally, it is necessary to detect small numbers of Nm in clinical specimens; bacterial loads in cerebrospinal fluid (CSF) of patients range from 3×10¹ to 3×10⁹ copies/mL.
4 × 10⁶ CFU/ml [8,9]. TaqMan rt-PCR has been shown to detect as few as 8 meningococcal genomes per reaction [4,5] and results are obtained within 2.5 hours.

ctrA may be the most frequently targeted gene to detect Nm using PCR [10]. However, the capsule locus, including ctrA, is subject to rearrangement [11–15], and 16% or more of carried meningococci have been shown to lack ctrA altogether [11,12]. Invasive NG meningococci can undergo similar rearrangements of the capsule region [J. Dolan Thomas, unpublished data], although these events may be less common than in carriage isolates.

The [Cu, Zn]-cofactored superoxide dismutase gene, sodC, is located 1.23 Mb from the capsule locus in the 2.27-Mb Nm serogroup B strain MC58 genome [16] and encodes the virulence factor Cu, Zn Sod. Cu, Zn Sod is a periplasmic enzyme [17], making it theoretically less susceptible to antigenic variation due to selective pressure than a cell-surface exposed molecule. sodC is believed to have been acquired by Nm via horizontal transfer from Haemophilus influenzae (Hi) [18]. There are no reports of meningococci that lack sodC, suggesting its importance to the survival of the organism in vivo and a strong selective pressure for its retention. However, sodC is not found in other Neisseria spp. [17–19].

The objective of this study was to improve detection of meningococci, especially of carriage isolates which may be ctrA-negative and NG, by developing a sensitive and specific rt-PCR assay for identification of all meningococci, regardless of capsule genotype or expression status.

Materials and Methods

Ethics statement

Ethics approval was not obtained from the CDC Institutional Review Board (IRB) for clinical specimens reported in this study because the specimens were sent to CDC for detection of meningitis etiology as part of reference lab functions. CDC IRB approval was not obtained for sodC testing of DNA extractions from Brazil and UK carriage specimens because those extractions were not tested by CDC researchers. DNA extractions of carriage specimens were not considered human specimens by the CDC, Emory University, and Children’s Hospital of Atlanta (CHOA) IRBs for this study, as they do not meet the definition of a living human subject. IRB approval was obtained by the institutions who collected the biological carriage specimens from the human subjects: (1) the National Ethics Research Committee and by the Regional Ethics Committee of the Hospital Materno Infantil, Secretary of Health of Goias State, Brazil; (2) the Emory University IRB; or (3) the National Health Service Research Ethics Committee (08/H1001/52), sponsored by the Royal Liverpool and Broadgreen University Hospitals Trust.

Bacterial strains and culture methods

Control isolates used for assay design and optimization are defined in Table S1.

626 cell lysates were used to determine the sensitivity of the sodC assay (Table 1 and Table S2), including lysates prepared from a temporally and geographically dispersed convenience sample of isolates from the CDC Meningitis Laboratory strain collection (received 1993–2006, n = 106) and all isolates from a US carriage study (n = 320) [20,21] known to be Nm by SAGS [1,2], rt-PCR serogrouping [5], NI strips (bioMérieux® sa), and Cystine Trypticase Agar (CTA) sugars (Remel) [1,2]. To further confirm identification, multilocus sequence typing (MLST) was performed on all U.S. carriage study and ctrA-negative NG isolates.

The specificity of the sodC assay for detecting only meningococci was determined using cell lysates from a total of 244 non-Nm isolates (Table 2).

Clinical specimens

CSF specimens from pediatric meningitis patients were cultured as soon as possible after collection. Specimens that were culture-negative were sent to CDC on ice for detection of meningitis etiology by the Marmara University School of Medicine in Istanbul, Turkey, and came from patients who met the case definition for purulent meningitis [leukocytosis (>100 cells/mm³) and either elevated protein (>100 mg/dl) or decreased glucose (<40 mg/dl)]. After DNA extraction and realtime PCR testing of all specimens for ctrA of Nm [5], lytA of S. pneumoniae [22], and bexA and/or boc2 of Hi, the subset of specimens chosen to test the sodC assay (n = 120) were either (1) positive for ctrA (n = 12) or (2) ctrA− lytA− bex/A− boc2− (n = 108).

37 U.S. clinical specimens were referred to CDC for detection or confirmation of bacterial meningitis etiology from January to June 2009, including CSF (n = 21), whole blood (n = 6), serum (n = 6), and tissues (n = 4) (Table 3).

Carriage specimens

The carriage specimens were obtained from three carriage studies. 1. In Goiania, Brazil, nasopharyngeal (NP) swabs (n = 223) were obtained from 154 children (ranging from 2–163 months of age) attending two daycare centers, 59 adult contacts of the attendees, and 10 daycare workers. The specimens were placed into skim milk-tryptone-glucose-glycerine (STGG) transport medium [23] and sent immediately to the Applied Microbiology Laboratory of Federal University of Goiás in Brazil for processing. The vials were then kept frozen during transport to CDC, where DNA extractions and rt-PCR were performed.

2. 291 posterior NP swab specimens were collected from a random sample of children 6 to 59 months of age who presented to the Emergency Department at CHOA at Egleston from March to August, 2009 [24]. Each swab specimen was immediately placed into 1 ml STGG [23]. Specimens were transported at room temperature to the clinical microbiology laboratory within 12 hours of collection for storage at −80°C until processing. An aliquot of the STGG from each specimen was transported on dry ice to the CDC Meningitis Laboratory, where DNA extractions and rt-PCR were performed.

3. A total of 33 NP swabs and 35 nasal washes (NWs) were taken from 24 participants ages 21–57 years during ≤7 visits in a Spring 2009 study conducted in the NIHR Biomedical Research Centre in Microbial Diseases at the Liverpool School of Tropical Medicine, Liverpool, United Kingdom (UK). Swab specimens were collected as previously described [25] with some modifications and placed directly into 1 ml STGG [23], then transported to the laboratory on wet ice for culture and processing. 900 μl of each specimen was frozen at −80°C for subsequent DNA extraction and rt-PCR in Liverpool.

NWs were collected as previously described [26] and the nasal fluid from both nostrils of each study participant was pooled. NWs were collected in this manner from each participant. Once at the lab, NW specimens were centrifuged at 1509 g for seven minutes and the pellet was re-suspended in 1 ml STGG. The processing of the NWs from this point was the same as that for the NP specimens.

DNA preparation and quantification

Genomic DNA was prepared for use in the various steps of assay design and optimization using the QIAamp DNA Mini
Kit (QIAGEN, Valencia, California) using Protocol C then quantified for use in standard curve experiments using a NanoDrop ND-1000 or 8000 spectrophotometer (Nanodrop Technologies, Wilmington, Delaware). Preparation of DNA from bacterial isolates was performed as previously described [5].
Gene sequencing

All extracted DNA was stored at −20°C and sequenced using the QIAsymphony SP System and the QIAsymphony Virus/Bacteria Midi Kit (QIAGEN Inc., UK) according to the manufacturer’s instructions. All extracted DNA was used for the sequencing of the sodC gene. The sodC consensus sequence was entered into Primer Express v.7 Program SeqMan.

rt-PCR primer and probe design

The sodC consensus sequence was entered into Primer Express 3.0 (Applied Biosystems). Primers and probes were designed for homology to other known sequences using the Basic Local Alignment Search Tool (BLAST) [27]. Primers were tested for optimal concentration in triplicate or quadruplicate in combination of final concentrations of 100, 300, 600, and 900 nM; the probe was tested in triplicate at final concentrations of 50, 100, 200, and 300 nM. The amplified product was located at nt 1427446 in MC58 (GenBank accession number AE002098.2).

rt-PCR

A Strategene Mx3005P (Agilent, La Jolla, California) and QuantiTect SYBR Green Master Mix (QIAGEN) were used to optimize primer concentrations. Cycle parameters were 2 minutes at 50°C, 10 minutes at 95°C, and then 50× (15 seconds at 95°C plus 1 minute at 60°C). Product dissociation curves were generated using one round of the following cycle parameters at the end of the primer optimization run: one minute at 95°C, 30 seconds at 55°C, and 30 seconds at 95°C. Master mixes contained 4.5 μl sterile PCR grade water (Roche Diagnostics), 12.5 μl TaqMan®2× PCR Master Mix (Applied Biosystems), 300 nM forward primer, 600 nM reverse primer, 100 nM probe, and 2 μl template DNA per total reaction volume of 25 μl. Each reaction plate that was run, cell lysates from known Nm served as positive external amplification controls, while no-template controls (NTCs) served as negative external amplification controls.

rt-PCR detection of ciaf in the MD isolates [20] was performed at the Laboratory of the Maryland Department of Health and Mental Hygiene using an Applied Biosystems 7500 or 7700 rt-PCR System (Applied Biosystems, Inc., Foster City, CA).

A Corbett Rotor-Gene 6000 (QIAGEN) and Corbett software 24 series 1.7 were employed in the UK for rt-PCR and data analysis on extractions of NWs and NP eluates. In every UK sodC reaction, 2.5 μl of extracted DNA was used.

Throughout this study, Ct values≤35 were considered positive; Ct values in the range of 36–40 equivocal; and Ct values>40 negative. Extracted DNA from equivocal clinical specimens was diluted 1:4 and 1:10 in an attempt to dilute possible inhibitors of the reaction. Dilutions were then re-tested in duplicate. If the average Ct of the diluted specimen fell below 35, that specimen was considered positive. If the average Ct of the diluted specimen remained in the 35–40 range, that specimen was considered negative.

MLST

MLST was performed as described by Maiden et al. [28]. Sequences were assembled and alleles were determined using Staden [29] v. 1.5.3 and Sequence Typing Analysis Retrieval System (STARS) v. 1.2a or the Meningococcus Genome Informatics Platform (MGIP) (http://mgip.biology.gatech.edu/) [30]. Sequence types (STs) and clonal complexes were assigned by

For CSF specimens and the Brazilian NP swab eluates, DNA extraction was conducted as previously described for clinical specimens [23]. DNA extractions were performed on the CHOA carriage study NP swab eluates using the MagNA Pure LC instrument and the DNA Isolation Kit III (Bacteria, Fungi) per the manufacturer’s instructions (Roche Diagnostics GmbH, Mannheim, Germany). DNA was extracted from the UK NP and NW specimens using the QIAsymphony SP System and the QIAsymphony Virus/Bacteria Midi Kit (QIAGEN Inc., UK) according to the manufacturer’s instructions. All extracted DNA was stored at −20°C.

Table 2. 244 non-Nm isolates were used to test the specificity of the sodC assay.

<table>
<thead>
<tr>
<th>Organism</th>
<th>n</th>
<th>sodC</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>M. catarrhalis</em></td>
<td>22</td>
<td>0</td>
</tr>
<tr>
<td><em>H. aphrophilus/paraphrophilus</em></td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td><em>H. aphrophilus</em></td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td><em>H. influenzae biogroup aegyptius</em></td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td><em>H. influenzae</em> serotype a</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td><em>H. influenzae</em> serotype b</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td><em>H. influenzae</em> serotype c</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td><em>H. influenzae</em> serotype d</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td><em>H. influenzae</em> serotype e</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td><em>H. influenzae</em> serotype f</td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td><em>H. influenzae</em> nontypeable (NTHi)</td>
<td>80</td>
<td>0</td>
</tr>
<tr>
<td><em>H. parainfluenzae</em></td>
<td>10</td>
<td>0</td>
</tr>
<tr>
<td><em>H. haemolyticus</em></td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td><em>N. lactamica</em></td>
<td>93</td>
<td>0</td>
</tr>
<tr>
<td><em>N. spp.</em></td>
<td>3</td>
<td>0</td>
</tr>
<tr>
<td><em>N. polysaccharaea</em></td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td><em>N. cinerea</em></td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td><em>N. subflava</em></td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td><em>N. sicca</em></td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td><em>N. gonorrhoeae</em></td>
<td>5</td>
<td>0</td>
</tr>
<tr>
<td><em>E. coli</em> K1</td>
<td>2</td>
<td>0</td>
</tr>
<tr>
<td><em>C. neoformans</em></td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td><em>S. aureus</em></td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td><em>S. pneumoniae</em></td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td><em>L. monocytogenes</em></td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td><em>A. pleuropneumoniae</em></td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td><em>S. choleraesuis</em></td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td><em>S. agalactiae</em></td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td><em>P. aeruginosa</em></td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td><em>C. diptheriae</em></td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td><em>B. pertussis</em></td>
<td>1</td>
<td>0</td>
</tr>
<tr>
<td>Total</td>
<td>244</td>
<td>0</td>
</tr>
</tbody>
</table>

192 N. lactamica, 4 N. gonorrhoeae, 19 M. catarrhalis, and 2 H. spp. from this panel were collected in a Georgia Nm carriage study [20]. 3 N. spp., 1 N. polysaccharaea, 2 M. catarrhalis, 9 H. parainfluenzae, 1 H. haemolyticus, 75 NTHi, and 1 Hie were collected in a Georgia Hib carriage study [24]. doi:10.1371/journal.pone.0019361.t002
shown). LLDs at a Ct value of 35 were found to be 39, 70, 101, and 82 genomes per reaction, yielding an average of 73 genomes detected per reaction. The average reaction efficiency was 100% and the average $R^2$ value was 0.9925.

Sensitivity of the sodC assay

Invasive (i.e., isolated from CSF, blood, joint fluid, or autopsy tissue, n = 76) and non-invasive (i.e., including carriage isolates and those from sputum or oral swab specimens, n = 30) known Nm isolates from the CDC Meningitis Laboratory Strain Collection were tested with the sodC assay. All isolates were positive for sodC, including 26 ctrA-negative NG isolates, with a median Ct value of 19.7, mean of 19.9±1.9, and range of 16.4 to 26.0 (Table 1 and Table S2). 518/520 (99.6%) of meningococcal carriage isolates from two U.S. carriage studies [20,24] were positive for sodC (median Ct, 16.9, mean 17.0±1.5, and range 13.5 to 29.3), while ctrA detected only 368/520 (70.8%) of these isolates (median Ct, 19.0, mean 19.2±2.7, and range 13.5 to 34.0) (Table 1 and Table S2). The two sodC-negative carried Nm were SASG NG, ctrA-negative but were confirmed to be Nm ST-1117 and ST-4788, both cc1117; both were isolated from the same study participant at different time points. Therefore, 176/178 (98.9%) SASG NG, ctrA-negative invasive and non-invasive Nm isolates were sodC-positive. Four sodC-negative carriage study isolates that were identified as Nm by conventional methods were re-investigated and shown to actually be non-Nm.

Specificity of the sodC assay

None of 35 non-Nm from various sources and none of 209 non-meningococcal carriage study [20,24] isolates were detected by sodC (100% specificity) (Table 2). Interestingly, sodC identified one isolate, M16160, as Nm when other standard carriage study tests could not correctly resolve its species. It was originally identified as querying http://pubmlst.org/neisseria or http://neisseria.org/nm/typing.

Nucleotide sequence accession numbers

The GenBank accession numbers for the Nm sodC sequences generated in this study are listed in Table S3.

Results

Primer and probe design

A consensus nucleotide sequence was built based on 2× coverage of sodC sequenced from each of 14 geographically and temporally dispersed groupable and NG meningococci (Table S1). Over the 439–473 out of 560 sodC nucleotides sequenced in these isolates, these meningococcal isolates were 99–100% identical to each other and 91% identical to an Hi sodC consensus that was built using two sequences from GenBank (accession numbers M84012 and AF549211). Given that sodC was likely acquired by Nm via horizontal transfer from Hi [18], primers and a probe that were chosen (Figure S1, Table 4). BLAST results showed that the primers had no homology over 78% nucleotide identity with any genes but meningococcal sodC. The only notable homology found for the probe was a two-nucleotide difference with H. parainfluenzae sodC; the primers, however, showed no homology to this gene.

Lower limit of detection (LLD)

Standard curves were generated by testing genomic DNA from four invasive meningococcal isolates with the sodC assay (data not shown). LLDs at a Ct value of 35 were found to be 39, 70, 101, and 82 genomes per reaction, yielding an average of 73 genomes detected per reaction. The average reaction efficiency was 100% and the average $R^2$ value was 0.9925.

Table 3. U.S. and Turkey clinical specimens tested for sodC and ctrA and compared to culture.

<table>
<thead>
<tr>
<th>Specimen</th>
<th>Total n</th>
<th>n sodC</th>
<th>% sodC</th>
<th>95% CI</th>
<th>n ctrA</th>
<th>% ctrA</th>
<th>95% CI</th>
<th>n Nm Culture Positive</th>
<th>% Culture Positive</th>
<th>95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>CSF</td>
<td>141</td>
<td>24</td>
<td>17.0</td>
<td>11% to 24%</td>
<td>20</td>
<td>14.2</td>
<td>9% to 21%</td>
<td>1</td>
<td>0.7</td>
<td>0% to 4%</td>
</tr>
<tr>
<td>Blood</td>
<td>6</td>
<td>1</td>
<td>16.7</td>
<td>0% to 64%</td>
<td>1</td>
<td>16.7</td>
<td>0% to 64%</td>
<td>0</td>
<td>0.0</td>
<td>0% to 46%</td>
</tr>
<tr>
<td>Serum</td>
<td>6</td>
<td>0</td>
<td>0.0</td>
<td>0% to 46%</td>
<td>0</td>
<td>0.0</td>
<td>0% to 46%</td>
<td>0</td>
<td>0.0</td>
<td>0% to 46%</td>
</tr>
<tr>
<td>Body Tissue</td>
<td>4</td>
<td>0</td>
<td>0.0</td>
<td>0% to 60%</td>
<td>0</td>
<td>0.0</td>
<td>0% to 60%</td>
<td>0</td>
<td>0.0</td>
<td>0% to 60%</td>
</tr>
<tr>
<td>Total</td>
<td>157</td>
<td>25</td>
<td>15.9</td>
<td>n/a</td>
<td>21</td>
<td>13.4</td>
<td>n/a</td>
<td>1</td>
<td>0.6</td>
<td>n/a</td>
</tr>
</tbody>
</table>

1Exact binomial 95% confidence interval.
2Culture was attempted for all 120 Turkey CSF specimens. With the exception of the one ctrA-negative, sodC-positive U.S. CSF specimen that was culture-positive, culture was either not attempted, not reported or, in one case, the isolate was nonviable two times for the other 36 U.S. clinical specimens.
3n/a, not applicable.

doi:10.1371/journal.pone.0019361.t003

Table 4. Primers and probe used in this study.

<table>
<thead>
<tr>
<th>Target Gene</th>
<th>Oligo Designation</th>
<th>Use</th>
<th>5’ to 3’ Nucleotide Sequence</th>
<th>Amplicon Size (bp)</th>
<th>Final Concentration (nM)</th>
</tr>
</thead>
<tbody>
<tr>
<td>sodC</td>
<td>Nm sodC-Fwd</td>
<td>PCR</td>
<td>sequencing</td>
<td>CCT TAT TAG CAC TAG CGG TTA G</td>
<td>400, 160</td>
</tr>
<tr>
<td></td>
<td>Nm sodC-Rev</td>
<td>PCR</td>
<td>sequencing</td>
<td>CCG GTC ATC TTT TAT GCT CCA A</td>
<td>537, 400, 160</td>
</tr>
<tr>
<td>sodC</td>
<td>Nm sodC Fwd 351</td>
<td>r-PCR</td>
<td>GCA CAC TTA GGT GAT TTA CCT GCA T</td>
<td>300</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Nm sodC Rev 478</td>
<td>r-PCR</td>
<td>CCA CCC GTG TGG ATC ATA ATA GA</td>
<td>127, 600</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Nm sodC Pb387</td>
<td>r-PCR</td>
<td>(FAM)-CAT GAT GGC ACA GCA ACA AAT CCT GTT T(BHQ1)</td>
<td>100</td>
<td></td>
</tr>
</tbody>
</table>

doi:10.1371/journal.pone.0019361.t004
N. polysaccharea/N. spp. by NH strip, but upon re-investigation, did not grow at room temperature on chocolate agar, indicating that it is not N. polysaccharea [1]. All 7 meningococcal housekeeping genes were readily amplified during MLST, suggesting that M16160 is indeed Nm (ST-7456, cc60), and again showing sodC to be a useful tool for definitive identification of carriage isolates.

Detection of Nm in clinical specimens

The ability of sodC to detect Nm was assessed and compared to ctrA as the target gene [5] using extracted DNA from 120 Turkish CSF specimens and 37 U.S. clinical specimens. ctrA detected Nm in 21/157 (13.4%) specimens (20 CSF and 1 blood), while sodC was positive for those 21 plus four additional CSF specimens (25/157, 15.9%) (Table 3 and Table S4). Therefore, sodC was 100% (95% confidence interval [CI]: 94–100%) sensitive compared to ctrA at detection of Nm from these clinical specimens, but specificity is difficult to calculate using ctrA as a reference test. The C values for the four ctrA-negative, sodC-positive CF extracts averaged 40.4 for ctrA while their sodC C1 values averaged 32.0.

One ctrA-negative, sodC-positive CF specimen was Nm culture-positive (1/157, 0.6%); the remaining specimens were either culture-negative (139/157) or culture was not attempted or not reported (17/157). Therefore, sodC was 88% (95% CI: 82–93%) specific compared to culture at detection of Nm from the clinical specimens for which culture was attempted (Table S4), but sensitivity is undetermined due to the low number of culture-positives.

Detection of Nm in carriage specimens

Results from all three carriage studies are summarized in Table 5. None (0/223, 0%) of the Brazilian NP swab eluate extractions tested at CDC were ctrA-positive while 3 (3/223, 1.3%) were sodC-positive, yielding C1 values of 29.1, 26.2, and 25.5. The ctrA-negative, sodC-positive specimens were negative for serogroups A, B, C, W135, X, and Y by PCR. In the 1/3 ctrA-negative, sodC-positive specimen that had a sufficient amount of extraction volume remaining, MLST confirmed the presence of meningococcal DNA (ST-923, cc198). All 23 Hi culture-positive Brazilian specimens were ctrA- and sodC-negative.

All (29/291, 100%) NP swab eluate extractions from the Georgia CHOIA carriage study were ctrA-negative and sodC-negative, as expected, since no Nm was cultured. These specimens were, however, culture-positive for N. spp. (n = 3), N. polysaccharea (n = 1), M. catarrhalis (n = 9), H. parainfluenzae (n = 9), H. haemolyticus (n = 1), and Hi (n = 76), further demonstrating the specificity of sodC.

S. aureus (n = 17), alpha-hemolytic streptococci (n = 18), M. catarrhalis (n = 5), diptheroids (n = 10), N. polysaccharea (n = 1), N. cinerea (n = 2), and N. meningitidis (n = 1) were cultured from the 68 UK carriage study specimens; the Nm culture-positive NW was ctrA-positive and sodC-positive. sodC and ctrA were negative for all of the non-Nm culture-positive specimens except 1 ctrA-positive, sodC-positive NW that grew N. cinerea and alpha-hemolytic streptococci. 1/33 (3%) NP swab eluates from the UK carriage study was ctrA-positive, 0/33 were sodC-positive, and 0/33 were Nm culture-positive. 3/33 (9.1%) NW extractions were ctrA-positive, 2/35 (5.7%) were also sodC-positive; of these, one (1/ 35, 2.9%) ctrA-positive, sodC-positive NW was Nm culture-positive. The ctrA-positive, sodC-negative NP specimen (average ctrA C1, 15.5 ± 0.3, average sodC C1, 37.3 ± 0.8) and the ctrA-positive, sodC-negative NW (average ctrA C1, value of 34.3 ± 0.2 and an average sodC C1 value of 33.1 ± 0.2) were both from patient 8, visit 3; this patient had the Nm-positive culture at visit 1 and the N. cinerea-positive culture at visit 2.

Discussion

Outbreaks of meningococcal disease occur in the U.S. and meningococcal disease continues to be a leading cause of bacterial meningitis worldwide. Regardless of location, when outbreaks of meningococcal disease occur or new vaccines are introduced, carriage studies are often performed to answer epidemiologic and biological questions (e.g., extent of the disease-causing clone circulating in the population) or demonstrate vaccine effects on carriage and therefore its potential impact on transmission and herd immunity.

Culture detection of Nm is 100% specific and should be attempted if possible, but it is limited by low sensitivity [31,32] and 24- to 72-hour incubation periods. The rapidity, sensitivity, and specificity of rt-PCR are rendering this technique an increasingly employed Nm detection method in clinical and reference laboratories [33–38]. Many laboratories use ctrA as a gene target for rt-PCR detection of Nm [3,5,10,35,38,39]. However, given that ctrA is not present in 16% or more of carriage isolates [11,12], it is not a suitable target gene for PCR on carriage study specimens. The reliability of ctrA-based PCR assays in detecting NG invasive Nm isolates [35] or specimens containing them, although rare, is also called into question by what is now known about the genetics of nongroupability in Nm carriage isolates [11,12] and the well-described genome plasticity of this naturally competent nasopharyngeal resident. Indeed, ctrA-based rt-PCR has recently been reported to generate false-negative results due to sequence variations in ctrA [40].

Table 5. Carriage specimens tested for sodC and ctrA and compared to culture.

<table>
<thead>
<tr>
<th>Specimen</th>
<th>Total n</th>
<th>n sodC</th>
<th>% sodC</th>
<th>95% CI</th>
<th>n ctrA</th>
<th>% ctrA</th>
<th>95% CI</th>
<th>n Nm Culture Positive</th>
<th>% Culture Positive</th>
<th>95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>NP swab eluate</td>
<td>547</td>
<td>3</td>
<td>0.5</td>
<td>0% to 2%</td>
<td>1</td>
<td>0.2</td>
<td>0% to 1%</td>
<td>0</td>
<td>0</td>
<td>0% to 1%</td>
</tr>
<tr>
<td>Nasal wash</td>
<td>35</td>
<td>2</td>
<td>5.7</td>
<td>0% to 19%</td>
<td>3</td>
<td>8.6</td>
<td>2% to 23%</td>
<td>1*</td>
<td>2.9</td>
<td>0% to 15%</td>
</tr>
<tr>
<td>Total</td>
<td>582</td>
<td>5</td>
<td>0.9</td>
<td>n/a</td>
<td>4</td>
<td>0.7</td>
<td>n/a</td>
<td>1*</td>
<td>0.2</td>
<td>n/a</td>
</tr>
</tbody>
</table>

1 Exact binomial 95% confidence interval.
2 This ctrA-positive NP swab eluate was sodC-negative.
3 Both of these sodC-positive NP swab eluates were ctrA-positive.
4 This Nm culture-positive NW was ctrA-positive, sodC-positive.
5 n/a, not applicable.
doi:10.1371/journal.pone.0019361.t005
The sodC assay was designed to detect Nm, especially among carriage isolates or specimens that lack ctrA, and it did, detecting 98.9% of carried and invasive ctrA-negative NG Nm isolates and from four ctrA-negative clinical and three ctrA-negative carriage specimens. sodC also surpassed culture in sensitivity by detecting Nm in 24 culture-negative clinical specimens and in at least one Nm culture-negative carriage specimen. The assay significantly improves specificity over culture, but sensitivity of sodC compared to culture is difficult to determine due to the lack of culture-positive clinical specimens received. This limitation underscores the benefit that the sodC real-time PCR assay will provide for the detection of Nm, especially in developing countries. Additionally, the specificity of sodC compared to ctrA cannot accurately be determined; the literature does not support the idea that ctrA is the ideal “gold standard” that is needed for a comparison of specificity [11,12,40]. sodC detects 73 genomes per 25-μl rt-PCR reaction, well within the range (≤10²–10⁷ CFU/ml) that would be found in CSF [9].

None of the 88 Hi isolates (Table 2) or the 100 Hi culture-positive Brazilian and CHOA carriage specimens tested were detected by this Nm sodC assay. sodC is also found in H. spp. other than Hi [41–43], but none of the 15 non-Hi H. spp. isolates (Table 2) and none of the 10 non-Hi H. spp. culture-positive CHOA carriage specimens tested were detected by sodC. To our knowledge, sodC has not been found in any other Neisseria spp. besides the meningococcus [17,18,44]. The species specificity of sodC was supported by the 100% specificity of the assay when tested on a panel of 244 non-Nm isolates, including 107 N. spp. isolates, and by 100% specificity of the assay when tested on carriage specimens, including 8 UK and CHOA specimens that were culture-positive for non-Nm N. spp.

The Ct value interpretation cutoffs and thermocycler parameters used for the sodC assay are the same as those used for detection of Nm serogroups A, B, C, W135, X, and Y [5] and for detection of Hi hpa [45] and S. pneumoniae lytA [23], making it convenient to perform multiple assays on isolates or specimens in one 96-well plate.

In summary, an extremely rapid, sensitive, and specific rt-PCR assay was developed and validated for the detection of meningococcal isolates and Nm from clinical and carriage specimens. This new diagnostic tool will be especially useful for confirming isolate species identification and detecting meningococci from carriage and clinical specimens, regardless of the organism’s encapsulation status.

Supporting Information

Figure S1  Nm sodC compared to Hi sodC and primer and probe location. sodC was sequenced from 14 groupable and NG meningococci and a consensus nucleotide sequence was generated. An H. influenzae sodC consensus was built using two sequences from GenBank (accession numbers M84012 and AF549211). Nucleotide differences from the meningococcal sequence are white letters highlighted in black in the Haemophilus sequence. The primers and probe (highlighted in gray) were chosen at positions not only containing at least three nucleotide differences between Nms and Hi, but also where sodC nucleotide sequence was conserved among the meningococcal isolates that were sequenced. The meningococcal sodC open reading frame is 560 nucleotides long; the portion of the gene depicted corresponds to nucleotides 253-513.

Table S1  Nm control isolates used for sodC assay design and optimization.

Table S2  626 Nm isolates used to test the sensitivity of the sodC assay.

Table S3  Nm sodC sequences generated in this study.

Table S4  Two-by-two contingency tables that were the basis for the calculation of sodC PCR sensitivity and specificity compared to culture and ctrA PCR. (a) Comparison of sodC PCR to culture for 140 clinical specimens. (b) Comparison of sodC PCR to ctrA PCR for 157 clinical specimens.

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Author Contributions

Conceived and designed the experiments: JDT CPH DAS MJT MCB KBL XZ RM SS JL-C APSdL JG. Performed the experiments: JDT CPH DAS MJT MCB KBL XZ RM SS JL-C APSdL JG. Analyzed the data: JDT CPH DAS MJT MCB KBL XZ RM SS KEA DSS LLH RAH ALA JL-C APSdL JG SG AS MB DS SJ SWS NEM LWM. Contributed reagents/materials/analysis tools: JDT CPH DAS MJT MCB KBL XZ RM SS KEA DSS LLH RAH ALA JL-C APSdL JG SG AS MB DS SJ SWS NEM LWM. Wrote the paper: JDT LWM. Assisted with editing, finalizing, & giving critical feedback on the scientific content of the manuscript: JDT CPH DAS MJT MCB KBL XZ RM SS KEA DSS LLH RAH ALA JL-C APSdL JG SG AS MB DS SJ SWS NEM LWM.


