Quality of metagenomic DNA extracted for molecular identification of microorganisms from CSF samples of patients with suspected cerebrospinal meningitis in northern Nigeria

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Abstract:

Background: Following an increase in the practice of starting antimicrobial therapy prior to clinical sample collection, the ability to confirm pathogenic microorganisms of bacterial meningitis has decreased by approximately 30%. Culture results may be false negative when fastidious or culture-resistant bacteria are involved or when patient samples are obtained after antimicrobial therapy has started. Molecular diagnosis using PCR can be performed directly on clinical samples after metagenomic DNA (mDNA) extraction not requiring live organisms for a positive result. The specific objectives of this study are to perform mDNA extraction directly from cerebrospinal fluids (CSF) using appropriate spin column method, and to determine the quality of the mDNA elute.

Methodology: Cerebrospinal fluid specimens were collected from 210 patients with suspected acute cerebrospinal meningitis (CSM) in the Federal Capital Territory and some States in Northern Nigeria during the 2017 and 2018 outbreak seasons. Metagenomic DNA was extracted from approximately 200µL of CSF specimens using the QiaGen QIAamp® DNA Mini kit specific for bacterial agents only. DNA quality check was performed on all DNA elutes using fluorometric, spectrophotometric and agarose gel electrophoresis methods.

Results: Of the 210 CSF samples analyzed microbiologically, Gram reaction was positive in 94 cases (44.8%) but only 17 (8.1 %) were culture positive for two of the three major bacterial causes of meningitis. One hundred and eighty (85.7%) samples had DNA concentrations ≥ 0.005 ng/µL, 55 (30.6 %) of these had DNA purity (A260/A280) of ≥ 1.7, 103 (57.2%) had purity value between 1.0 - 1.69, 14 (7.8%) had value of 0.57 - 0.99, and 8 (4.4%) failed purity evaluation with value of 0.00 at A260/A280.

Conclusion: The essence of mDNA extraction is multipurpose. A multiplex PCR can be performed on the extracted mDNA to interrogate the presence of microbial pathogens of interest using specific primers and probes (when applicable). Quality mDNA from CSF samples will ensure successful qPCR results for rapid and accurate detection of bacterial pathogens in meningitis. This will eliminate the challenges associated with traditional culture methods.

Keywords: Meningitis, CSF, DNA Quality Check, Fluorometry.

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Qualité de l’ADN métaganomique extrait pour l’identification moléculaire des microorganismes à partir d’échantillons de LCR de patients suspectés de méningite cérébrospinale dans le nord du Nigéria

*1,3Peletiri, I. C., 1Ikeh, E. I., 2Nna, E., 3Ndike, U. P., 4Usman, Y. B., 5Durfa, L. D., 6Okonkwo, C. N., 7Murtala, R., et 2Nnajide, C. R.
Abstrait:

Contexte: Suite à une augmentation de la pratique de commencer un traitement antimicrobien avant le prélèvement d’échantillons cliniques, la capacité à confirmer les microorganismes pathogènes de la méningite bactérienne a diminué d’environ 30%. Les résultats de la culture peuvent être faux négatifs lorsque des bactéries exigeantes ou résistantes à la culture sont impliquées ou lorsque des échantillons de patients sont prélevés après le début du traitement antimicrobien. Le diagnostic moléculaire par PCR peut être réalisé directement sur des échantillons cliniques après extraction d’ADNm métagénomique (ADNm) ne nécessitant pas d’organismes vivants pour un résultat positif. Les objectifs spécifiques de cette étude sont d’effectuer l’extraction de l’ADNm directement à partir de fluides céphalo-rachidiens (LCR) en utilisant de la colonne de rotation appropriée, et de déterminer la qualité de l’élué d’ADNm.


Résultats: Sur les 210 échantillons de LCR analysés microbiologiquement, la réaction de Gram était positive dans 94 cas (44,8%), mais seulement 17 (8,1%) étaient positives en culture pour deux des trois principales causes bactériennes de la méningite. Cent quatre-vingt (85,7%) échantillons avaient des concentrations d’ADN ≥ 0,005 ng/µL, 55 (30,6%) d’entre eux avaient une pureté d’ADN (A260/A280) ≥ 1,7, 103 (57,2%) avaient une valeur de pureté comprise entre 1,0 et 1,69, 14 (7,8%) avaient une valeur de 0,57 à 0,99, et 8 (4,4%) ont échoué l’évaluation de la pureté avec une valeur de 0,00 à A260/A280.

Conclusion: L’essence de l’extraction d’ADNm est polyvalente. Une PCR multiplex peut être effectuée sur l’ADNm extrait pour interroger la présence d’agents pathogènes microbiens d’intérêt en utilisant des amorces et des sondes spécifiques (le cas échéant). Un ADNm de qualité provenant d’échantillons de LCR assurera des résultats de qPCR réussis pour une détection rapide et précise des bactéries pathogènes dans la méningite. Cela éliminera les défis associés aux méthodes de culture traditionnelles.

Mots clés: méningite, LCR, contrôle de la qualité de l’ADN, fluorométrie

Introduction:

The continuous yearly outbreak of acute meningitis over the years no doubt had left behind very sad memories, moments in the minds and life of individuals (sufferers), family members, friends, communities and nations alike especially in the Meningitis Belt of Africa. Meningitis has been reported as one of the deadliest diseases that has been plaguing West Africa for decades. The sub-Saharan Africa was been plagued by large epidemics of meningo-coccal meningitis for a century, leading to it being labelled the ‘meningitis belt’ (1), spanning 26 countries (Fig 1).

Epidemics usually occur in the dry season which commences from December to June, with an epidemic wave that can last two to three years but dies out during the inter-

vening raining seasons (2). In 1998, the World Health Organization attributed several factors to be associated with the development of epide-

mics in the Africa’s meningitis belt. These factors include medical conditions (immunological susceptibility of the population), demo-

graphic conditions (travel and large population displace-

ments), socioeconomic conditions (overcrowding, poor hygiene and living conditions), climatic conditions (drought and dust storms), and concurrent acute respiratory infections (3). In Nigeria, about 5000 cases of mening-

itis occur every year with loss of many lives (4). The Nigeria Centre for Disease Control (NCDC) and Federal Ministry of Health (FMoH) Weekly Epidemiological Report from 2012 to 2018 revealed that this problem persists. Suspected cerebrospinal meningitis (CSM) cases within these seven years were 21,353; of which only
643 cases (3.01 %) were laboratory confirmed. The number of deaths (case fatality ratio, CFR) was 1,347 (6.31 %) (5).

Bacterial meningitis remains a serious global health problem as well as a life-threatening condition that requires prompt recognition and treatment. It is also documented that, over 1.2 million cases of bacterial meningitis are estimated to occur worldwide each year (3). Without treatment, the case-fatality rates vary from 10% to greater than 50% (6), and can be as high as 70%, with one in five (20%) survivors left with permanent sequelae including hearing loss, neurologic disability, or loss of a limb (7).

The diagnosis of bacterial meningitis rests heavily on examination of CSF collected through lumber puncture (8). The presumptive identification of Neisseria meningitidis, Streptococcus pneumoniae and Haemophilus influenzae as well as other bacteria can be made on the basis of cytological examination of the CSF, specific colony morphology on blood and/or chocolate agar, staining properties on Gram stain or by detection of specific antigens in the CSF by latex agglutination test or a rapid diagnostic test (9). Although culture technique has been recommended as the ‘gold standard’ test because cultured bacteria are sources of data for antibiotic susceptibility, complete sub-typing, expression of antigens that are to be included in future vaccines, and understanding the pathophysiology of isolates, specimens that do not yield any culture growth can still be analyzed by molecular methods using metagenomic DNA (mDNA) extracted from clinical samples (9). A further probing into this statement by the CDC (9), revealed that they were actually referring to “metagenomic protocol” (either consciously or otherwise) aimed at tackling the ‘yielded no growth syndrome’ or abysmal low yield of CSF culture results. Culture was referred to as the ‘gold standard’ before now because it was the only method that provided evidence for the presence of any aetiologial agent which can be used for further down-stream activities. In low income resource countries that do not have molecular diagnostic methods such as PCR, CSF culture method remains the best option.

Metagenomics is being described as the direct study of genetic materials recovered or extracted from microbial communities present in environmental samples, which take advantage of the rich diversity of genes and biochemical reactions of millions of non-cultivated and unculturized microorganisms (10). Metagenomics has been in the practice of microbiology for a while. Far away in 1935, Henrici and Johnson reported the known age long standard methods in bacteriology of “pure culture isolation and observation upon artificial media which often yield only an incomplete knowledge of a particular microbial flora” (11). Truly, microbial cultures have always been used to determine the microbial composition, but at the present, reports state that a large proportion of microorganisms in each ecosystem cannot be cultured with traditional tools, and their detection is only possible with DNA sequencing of their genetic fingerprints, the so-called metagenome (12). However, it should be clear at this point that the detection of bacteria in clinical or environmental samples by way of metagenomic approach is not only possible with DNA sequencing, but with molecular detection using multiplex PCR protocol with specific primers and probes (when applicable). Reports on estimates have it that cultured microorganisms account for less than 20% of the real phylogenetic diversity of pro-
karyotes (13). Bacteria may be recalcitrant to culturing for diverse reasons such as lack of necessary symbionts, nutrients, or surfaces; excess inhibitory compounds; incorrect combinations of temperature, pressure, or atmospheric gas composition; accumulation of toxic waste products from their own metabolism; and intrinsically slow growth rate or rapid dispersion from colonies (14).

Due to the increase in the practice of starting antimicrobial therapy prior to clinical sample collection (15-18), the confirmation of aetiological agents of bacterial meningitis has been reported to decrease by about 30% (16, 17). The use of molecular diagnostic protocol (especially PCR analysis) offers the advantages of detecting the DNA of serogroup-specific N. meningitidis as well as other implicating microorganisms, and not requiring live organisms for a positive result (19). In 2012, Foxman reported that advanced molecular techniques have provided the opportunity to detect trace amounts of genetic materials of a pathogen in various specimens with sensitivity that is far beyond culture-based methods (20).

Molecular techniques especially PCR-based assay, have become available to provide an early and accurate diagnosis of bacterial meningitis (21). This assay can detect as few as 10-100 CFU/mL of bacteria in CSF (22). PCR can be performed directly on clinical samples; the viability or otherwise of any organism present does not affect the result, this being that DNA can be extracted from clinical samples (typically blood and CSF) (9). The use of molecular assays for detection of aetiological agents of meningitis directly from CSF is now an established protocol.

The efficient extraction/preparation of DNA template is a necessary step for any real-time PCR (9) to meet the required DNA quality in terms of concentration, purity and quantity. The goal of DNA extraction is to lyze the bacterial cells in the specimens to maximize bacterial DNA yield and quality while removing any PCR inhibitors (i.e. salts, proteins), and dissolve the DNA in a buffer compatible with the enzymes used in the next analytical step while concentrating the extracted DNA at the same time (9).

Commercial DNA extraction kits are available for culture, blood and body fluids (CSF inclusive). One of such DNA extraction kit is the Qiagen QIAamp(R) DNA Mini Kit which provides fast and easy method for purification of total genomic DNA for reliable PCR, and can purify total DNA (e.g. metagenomic) from whole blood, plasma, serum, buffy coat, bone marrow, CSF, lymphocytes, cultured cells, tissue, and forensic specimens (23). The Qiagen QIAamp DNA Mini kit is a spin column technology in which DNA is selectively absorbed onto silica membrane (24). DNA purified using QIAamp kits is up to 50 kb in size, with fragments of approximately 20-30kb predominating. DNA of this length denatures completely during thermal cycling and can be amplified with high efficiency.

It is mandatory to determine the quality (concentration and purity) of extracted metagenomic DNA before use for PCR assays. DNA yield can be assessed using various methods including absorbance (optical density), agarose gel electrophoresis, or fluorescent DNA-binding dyes. The Eppendorf BioPhotometer Plus instrument used for quality check measures optical density (absorbance values) at 230, 260 and 280 nanometres and converts optical densities to concentrations. Dependent on the method, the results can be calculated through fixed factors, standards, or curve calibration. In addition to the results, the device also displays the absorbance values and some other important details, such as the common absorbance quotients e.g. A260/A280 and A230/A280 ratio for nucleic acid calculations. Purity is determined by calculating the ratio of absorbance at 260 nm to absorbance at 280 nm. Pure DNA (good quality DNA) has A260/A280 ratio of 1.7–1.9, although reading lower than 1.6 does not render the DNA unsuitable for any application, but lower ratios may indicate the presence of more contaminants (25). Elution buffer for genomic DNA is usually 1 x Tris EDTA buffer at pH 8.0.

For checking DNA quality and yield, RNase/DNase free water (nuclease free water) is used to dilute samples and to zero the spectrophotometer while the absorbance is measured at A260 and A280 nm. Both DNA and RNA are measured with spectrophotometer, however, to measure only DNA, a fluorometer must be used (23). Quality check of DNA extract by agarose gel electrophoresis is one of the most frequently used techniques in life sciences (26). DNA fragments loaded on agarose gels would have been stained with ethidium bromide and detected by an ultraviolet (UV) transilluminator system (27). Agarose gel electrophoresis is also another way to quickly estimate DNA concentration.

Fluorescence measurement of nucleic acids is based on the use of fluorogenic dyes that bind selectively to DNA or RNA. Dyes only emit signal when bound to the target, and signal is measured by fluorometers. Sample is excited with filtered light (at the excitation wavelength), and the emitted light (at the emission wavelength) is recorded by a detector (28). The objective of this study is to evaluate the quality of mDNA (concentration, purity, and amount) extracted by spin column technique (Qiagen
Metagenomic DNA extraction from CSF samples

Materials and method:

Study settings
The study sites were in Federal Capital Territory (FCT): National Hospital, Abuja; all District/General Hospitals in the FCT (Asokoro, Wuse, Maitama, Garki, Gwarinpa, Bwari, Kubwa, Kuje, and Nyanya), and some States in Northern Nigeria. However, CSF samples were not received from any District/General Hospital in the FCT; but from National Hospital, Abuja and also from Kebbi, Plateau, Sokoto, and Zamfara States in Northern Nigeria during outbreak seasons of February – May 2017 and January – April, 2018.

Ethical consideration
Ethical approvals were obtained from the Ethics Committees of National Hospital, Abuja (NHA/EC/034/2015), Federal Capital Development Authority Health Services (FHREC/2017/01/27/03-04-17), Kebbi State Ministry of Health (MOH/KSREC/VOL.1/56/No 101.3/2015), Plateau State Ministry of Health (MOH/MIS/202/VOL.T/X,2017), Sokoto State Ministry of Health (SMH/1580/V.IV, 2017), and Zamfara State Ministry of Health (ZSHREC/02/03/2017). A letter of introduction from the Nigeria Centre for Disease Control (NCDC) of the Federal Ministry of Health (Ref. MH/2768/S.162/III) was obtained to cover for all outbreak sites in the country. Written informed consent for storage and future use of unused samples, sample materials and data transfer agreement, were also obtained.

Subjects
All hospitalized patients (all ages and gender) with clinical symptoms of meningitis as reviewed by the attending physicians were included in the study. Patients who did not give informed consent and sites that did not grant approval were excluded from the study.

Sample size and sampling method
The sample size was determined using the Cochran formula (29) for calculating simple proportion; 

\[ n_s = \frac{z^2 pq}{e^2} \]

where \( n_s \) is the minimum required sample size, ‘\( z \)’ is the selected critical value of desired confidence level at 95% (standard value of 1.96), \( 'p' \) is the estimated proportion of an attribute that is present in the population [estimated prevalence of meningitis in Zamfara State of 13.7% (30)], \( 'q' \) is 1-p and \( 'e' \) is the desired level of precision (margin of error at 5%; standard value of 0.05). Therefore, the estimated sample size was 181.7 which was adjusted to 210 samples after calculating for 10% attrition. The subjects were recruited consecutively until the sample size was attained.

Collection and transportation of CSF specimens
CSF was collected into sterile containers by experienced physicians after performing lumbar puncture under aseptic conditions. The samples were transported to the laboratory at the various sites and kept at -20°C before being transported in ice-packs to Abuja for onward transfer to the Safety Molecular Pathology Laboratory, Enugu, where the samples were kept at -80°C until DNA extraction was carried out.

Metagenomic DNA extraction
The QiaGen QIAamp(R) DNA Mini kit was used for DNA extraction of the CSF samples, bacterial isolates and three ATCC control strains (\( N. meningoitedes \) serogroup B ATCC 13090, \( H. influenzae \) Type B, Biotype 1 ATCC 10211, and \( S. pneumoniae \) serotype 19F ATCC 49619). Approximately 200μL each of the CSF, bacterial isolates from the CSF (kept in 10% Skim milk with 15% glycerol) and ATCC bacterial control strains in TE buffer, were transferred into 2.0mL microcentrifuge (Eppendorf) tubes. 20μL of proteinase K was added to all samples, vortexed at 2000 rpm for 5 seconds, and incubated at 56°C for 15 minutes. 60μL buffer AL was added, mixed thoroughly by vortexing for 15 seconds, incubated at 70°C for 10 minutes and briefly centrifuged to remove drops from the lid of the tube. 200μL ethanol (96–100%) was added, vortexed for 15 seconds and briefly centrifuged to remove drops from the lid of the tube.

The mixture was pipetted onto the QIAamp Mini spin column (in a 2ml collection tube), centrifuged at 6000xg (8000rpm) for 1 min, and the flow-through and collection tube discarded. The QIAamp Mini spin columns were placed in a new 2ml collection tube, 500μL buffer AW1 was added, centrifuged at 6000xg (8000rpm) for 1 min and the flow-through and collection tube discarded. The QIAamp Mini spin columns were placed in a new 2ml collection tube, 500μL buffer AW2 was added, centrifuged at full speed of 20,000 x g (14,000 rpm) for 3 min and the flow-through and collection tube discarded. The QIAamp Mini spin columns were then placed in a new 1.5 ml collection tube and 60μL buffer AE added, incubated at room temperature for 1min and centrifuged at 6000xg (8000rpm) for 1min to elute the mDNA. Eluted DNA samples were labelled accordingly for quality check and appropriately stored for later use.
**Metagenomic DNA extract quality check**

The quality of the extracted mDNA was performed by the fluorometric method (for concentration) and spectrophotometric method (for purity). The agarose gel electrophoresis method was used to access few DNA samples for the presence or absence of bands only.

**Fluorescence method for DNA concentration**

The Qubit 3.0 fluorometer instrument (Invitrogen Life Technologies, now Thermo-Fisher) was used to determine the concentration of the extracted mDNA from the samples. Assay components were equilibrated at room temperature; the Qubit(R) working solution was prepared by diluting Quant - IT™ dsDNA HS reagent 1:200 in Quant - IT™ dsDNA HS buffer. 200µl of working solution was prepared for each standard and sample. The assay tubes were prepared according to Table 1.

All tubes were vortexed for 2–3 sec. The tubes were incubated for 2 minutes at room temperature. The tubes were inserted in the Qubit 3.0 fluorometer and readings taken. The Qubit 3.0 fluorometer was calibrated using the readings of the Standard Assay Tubes (8 in number) with concentrations range of 0.0 ng/µl to 10.0ng/µl, and prepared a standard curve to determine DNA amounts in user samples (unknown DNA sample concentrations). For the calibration curve, data from Qubit 3.0 were entered into GraphPad Prism and linear regression of DNA standards was determined (Table 2) and used in reading the relative fluorescence unit of samples (Table 3).

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**Table 1: Protocol for preparing assay tubes for fluorometric method**

<table>
<thead>
<tr>
<th>Volume of solution/analyte</th>
<th>Standard Assay Tubes</th>
<th>Unknown DNA (user) samples Assay Tubes</th>
</tr>
</thead>
<tbody>
<tr>
<td>Volume of working solution (from step 2) to add</td>
<td>190 µl</td>
<td>195 µl</td>
</tr>
<tr>
<td>Volume of standard (from kit) to add</td>
<td>10 µl</td>
<td>-</td>
</tr>
<tr>
<td>Volume of user sample to add</td>
<td>-</td>
<td>5 µl</td>
</tr>
<tr>
<td>Total volume in each Assay Tube</td>
<td>200 µl</td>
<td>200 µl</td>
</tr>
</tbody>
</table>

Thin–walled, clear 0.5ml PCR tubes were used. Acceptable tubes include Qubit(R) assay tubes (set of 500 – Cat No. Q32856) or Axygen PCR – 05 – C tubes (VWR, Part No. 10011 - 830). The minimum assay volume must be 200 µl.

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**Table 2: Characteristics of the Linear Regression for DNA check by Fluorometric Method (Qubit 3.0)**

<table>
<thead>
<tr>
<th>Best-fit values</th>
<th>Relative Fluorescent Unit (RFU)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Slope 1651 ± 53.21</td>
<td>487.1 ± 268.9</td>
</tr>
<tr>
<td>Y-intercept when X= 0.0</td>
<td>- 0.2950</td>
</tr>
<tr>
<td>X-intercept when Y= 0.0</td>
<td>0.0006056</td>
</tr>
<tr>
<td>1/slope</td>
<td>0.0006056</td>
</tr>
<tr>
<td>95% Confidence Intervals Slope</td>
<td>1503 to 1799</td>
</tr>
<tr>
<td>Y-intercept when X= 0.0</td>
<td>-259.4 to 1234</td>
</tr>
<tr>
<td>X-intercept when Y= 0.0</td>
<td>-0.8004 to 0.1478</td>
</tr>
<tr>
<td>Goodness of Fit</td>
<td>R square 0.9959</td>
</tr>
<tr>
<td>Sy.x</td>
<td>464.5</td>
</tr>
<tr>
<td>Is slope significantly non-zero?</td>
<td>F 962.9</td>
</tr>
<tr>
<td>DFn, DFD</td>
<td>1.000, 4.000</td>
</tr>
<tr>
<td>P value</td>
<td>&lt; 0.0001</td>
</tr>
<tr>
<td>Deviation from zero?</td>
<td>Significant</td>
</tr>
<tr>
<td>Data</td>
<td>Number of X values 6</td>
</tr>
<tr>
<td>Maximum number of Y replicates</td>
<td>1</td>
</tr>
<tr>
<td>Total number of values</td>
<td>6</td>
</tr>
<tr>
<td>Number of missing values</td>
<td>0</td>
</tr>
</tbody>
</table>
Table 3: Fluorometric readings of standards

<table>
<thead>
<tr>
<th>Concentration (ng/µl)</th>
<th>RFU value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Std1 10.00</td>
<td>17203.03</td>
</tr>
<tr>
<td>Std2 8.00</td>
<td>9922.97</td>
</tr>
<tr>
<td>Std3 6.00</td>
<td>9735.85</td>
</tr>
<tr>
<td>Std4 4.00</td>
<td>7538.76</td>
</tr>
<tr>
<td>Std5 1.00</td>
<td>2423.75</td>
</tr>
<tr>
<td>Std6 0.50</td>
<td>989.15</td>
</tr>
<tr>
<td>Std7 0.00</td>
<td>532.96</td>
</tr>
</tbody>
</table>

Table 4: Spectrophotometer Method – Operation

<table>
<thead>
<tr>
<th>Method group</th>
<th>Method</th>
<th>Description</th>
<th>Wavelength</th>
</tr>
</thead>
<tbody>
<tr>
<td>DNA</td>
<td>dsDNA</td>
<td>Calculating the concentration of DNA with evaluation via factor. Already pre-programmed factor ex-factory</td>
<td>Measuring wavelength: 260nm, Secondary wavelength to check for purity: 280nm</td>
</tr>
</tbody>
</table>

Spectrophotometric method for DNA purity

In the spectrophotometric method, the Eppendorf BioPhotometer Plus instrument was used to determine the purity of the extracted metagenomic DNA from the samples. Measuring procedure was according to literature insert in the manufacturer’s manual, with the instrument set for dsDNA and sample dilutions of 5µL sample + 95µL diluent for reading at A₂₆₀/A₂₈₀. The Eppendorf BioPhotometer Plus instrument switched-on to initialize. Sample preparation (95µL of diluent distilled water) was pipetted into appropriately labelled tubes. 5µL of mDNA extract was added to the corresponding labelled tubes. The sample was transferred into a clean cuvette shaft of outside diameter 12.5mm x 12.5mm. The instrument was set at blank (zero) before reading at A₂₆₀/A₂₈₀ wavelength (Table 4). Result was recorded for purity value at A₂₆₀/A₂₈₀ and dsDNA concentration in µg/mL but readings of the dsDNA concentration were disregarded because of inconsistent values (non-reproducibility of readings).

Gel electrophoresis method

In the gel electrophoresis method, the Biorad Horizontal Gel Electrophoresis tank was used in running of some samples. The set gel (1.5% agarose) was well placed into the tank and filled to the brim with 0.5xTris Borex EDTA (TBE). 5µL of mDNA extract was pipetted into microlitre plate wells appropriately. 3µL of loading dye (10 x Dream Taq Green Buffer which includes 20mM MgCl₂) was added into all samples in the microlitre plate wells and well mixed. The loading dye helps the DNA extract (sample) to sink into the well of the gel. Each gel well was loaded with the sample and covered with adequate 0.5xTBE. The electrophoresis tank is connected to the power pack with positive and negative terminals, and switched on. The gel was run at 100 volts for 45–60 min. The gel was then transferred to the UV trans-illuminator that is fitted with a camera system which is viewed on a computer connected to the camera system. Using the GenoSpot programme and EOS Utility (for the Camera) that are installed on the computer, snap shot gel pictures of the mDNA bands were taken, saved and labelled appropriately.

Calculating amount of mDNA extracts

The mDNA concentration (ng/µL) was measured by the Qubit Fluorometer 3.0g. The DNA yield = DNA concentration x eluted volume (60µL) per 200µL of CSF. The amount of DNA = DNA concentration x 5µL per qPCR reaction.

Results:

Of the 210 subjects recruited into the study, 129 (61.4%) were males, comprising 104 (49.5%) children (<15 years of age) and 25 (12%) adults while females were 81 (38.6%), comprising 66 (31.4%) children (<15 years) and 15 (7.1%) adults (Table 5). Following microbiological analysis, Gram reaction was positive in 94 (44.8%) samples while only 17 (8.1%) were culture positive for two of the three bacteria under study (Table 6).

Table 7 shows the summary of results obtained from linear regression for mDNA concentration by fluorometric method (Qubit 3.0) on the 210 CSF samples. Metagenomic DNA was extracted from 180 (85.7%) samples with concentrations of ≥ 0.005 ng/µL (relative fluorescent unit (RFU) value of 537.32) being the lowest limit of detection (LOD), while 30 (14.3%) had DNA concentration less than 0.005 ng/µL.
ng/µl. The amounts of mDNA present in the 180 (85.7%) samples were DNA concentration of 0.03–50.5ng/µl, DNA yield of 1.8–3030 µg and DNA amount of 0.15–252.5 ng/µl. Table 8 is the summary results of mDNA purity values at \(A_{260}/A_{280}\) and concentration of ≥ 0.005 ng/µl.

### Table 5: Age group and gender distribution of subjects recruited for the study

<table>
<thead>
<tr>
<th>Age group (years)</th>
<th>Male (%)</th>
<th>Female (%)</th>
<th>Total (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt; 15</td>
<td>104 (49.5)</td>
<td>66 (31.4)</td>
<td>170 (81.0)</td>
</tr>
<tr>
<td>&gt; 15</td>
<td>25 (11.9)</td>
<td>15 (7.1)</td>
<td>40 (19.0)</td>
</tr>
<tr>
<td>Total</td>
<td>129 (61.4)</td>
<td>81 (38.6)</td>
<td>210 (100)</td>
</tr>
</tbody>
</table>

### Table 6: Results of Gram reaction, culture and metagenomic DNA on the CSF samples

<table>
<thead>
<tr>
<th>Test</th>
<th>CSF samples</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gram reaction</td>
<td>94</td>
</tr>
<tr>
<td>Culture</td>
<td>17</td>
</tr>
<tr>
<td>Metagenomic DNA (≥0.005ng/ul)</td>
<td>180</td>
</tr>
</tbody>
</table>

CSF = cerebrospinal fluid

### Table 7: Linear regression for mDNA concentration by fluorometric method (Qubit 3.0) on CSF samples

<table>
<thead>
<tr>
<th>Concentration (ng/ul)</th>
<th>RFU value</th>
<th>No of samples</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.005 - ≥ 10.00</td>
<td>537.32 - ≥ 17203.03</td>
<td>180</td>
<td>85.7</td>
</tr>
<tr>
<td>0.00 - &lt; 0.005</td>
<td>532.96 - &lt; 537.32</td>
<td>30</td>
<td>14.3</td>
</tr>
<tr>
<td>Total</td>
<td>210</td>
<td></td>
<td>100</td>
</tr>
</tbody>
</table>

RFU = Relative Fluorescent Unit; mDNA = metagenomic DNA

### Table 8: Spectrophotometric results of mDNA purity at \(A_{260}/A_{280}\) and concentration of ≥ 0.005 ng/µL

<table>
<thead>
<tr>
<th>Purity @ (A_{260}/A_{280})</th>
<th>Number of samples</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>≥ 1.7</td>
<td>55</td>
<td>30.6</td>
</tr>
<tr>
<td>1.0 – 1.69</td>
<td>103</td>
<td>57.2</td>
</tr>
<tr>
<td>0.57 – 0.99</td>
<td>14</td>
<td>7.8</td>
</tr>
<tr>
<td>0.00</td>
<td>8</td>
<td>4.4</td>
</tr>
<tr>
<td>Total</td>
<td>180</td>
<td>100</td>
</tr>
</tbody>
</table>
Fig 2 shows the linear regression plot for mDNA concentration. Fig 3 shows gel picture of ATCC bacterial control strains that had DNA purity at $A_{260}/A_{280}$ of 1.53 ($N. meningitidis$), 1.48 ($H. influenzae$), and 1.57 ($S. pneumoniae$) and Fig 4 shows the gel electrophoresis picture of mDNA of samples that had purity range of 1.0–4.37 at $A_{260}/A_{280}$.

Fig 2: Linear Regression Plot for DNA Concentration

Lane 1 – DNA ladder; Lane 2 – $N. meningitidis$ (DNA purity @ $A_{260}/A_{280}$ – 1.53); Lane 3 – $H. influenzae$ (DNA purity @ $A_{260}/A_{280}$ – 1.48); Lane 4 – $S. pneumoniae$ (DNA purity @ $A_{260}/A_{280}$ – 1.57); Lane 5 – $N. meningitidis$ (DNA purity @ $A_{260}/A_{280}$ – 1.53); Lane 6 – $H. influenzae$ (DNA purity @ $A_{260}/A_{280}$ – 1.48)

Fig 3: Gel electrophoresis picture of ATCC bacterial control strains

Lane 1: DNA Ladder (50 bp); Lane 2: IP01N (DNA purity @ $A_{260}/A_{280}$ – 1.23; DNA Conc. 1.3 ng/µL); Lane 3: IP07N (DNA purity @ $A_{260}/A_{280}$ – 0.95; DNA Conc. 10.8 ng/µL); Lane 4: IP026N (DNA purity @ $A_{260}/A_{280}$ – 1.3; DNA Conc. 0.14 ng/µL); Lane 5: IP35N (DNA purity @ $A_{260}/A_{280}$ – 3.1; DNA Conc. 0.04 ng/µL); Lane 6: IP62N (DNA purity @ $A_{260}/A_{280}$ – 1.9; DNA Conc. 0.14 ng/µL); Lane 7: IP95S (DNA purity @ $A_{260}/A_{280}$ – 1.0; DNA Conc. 5.9 ng/µL); Lane 8: IP101N (DNA purity @ $A_{260}/A_{280}$ – 1.2; DNA Conc. 0.12 ng/µL); Lane 9: IP128H (DNA purity @ $A_{260}/A_{280}$ – 1.1; DNA Conc. 0.97 ng/µL); Lane 10: IP147N (DNA purity @ $A_{260}/A_{280}$ – 1.2; DNA Conc. 5.8 ng/µL); Lane 11: IP157H (DNA purity @ $A_{260}/A_{280}$ – 8.1; DNA Conc. 6.1 ng/µL); Lane 12: IP178S (DNA purity @ $A_{260}/A_{280}$ – 1.4; DNA Conc. 1.8 ng/µL); Lane 13: IP189N (DNA purity @ $A_{260}/A_{280}$ – 1.8; DNA Conc. 4.7 ng/µL); Lane 14: IP209N (DNA purity @ $A_{260}/A_{280}$ – 2.0; DNA Conc. 1.1 ng/µL); Lane 15: Nm ATCC (DNA purity @ $A_{260}/A_{280}$ – 1.5; DNA Conc. 15.1 ng/µL); Lane 16: Hi ATCC (DNA purity @ $A_{260}/A_{280}$ – 1.48; DNA Conc. 20.0 ng/µL); Lane 17: Sp ATCC (DNA purity @ $A_{260}/A_{280}$ – 1.6; DNA Conc. 16.6 ng/µL)

Fig 4: Gel electrophoresis picture of mDNA samples (DNA purity @ $A_{260}/A_{280}$ of 1.10 – 4.37)
Discussion:

The findings of our study showed that Gram reaction was positive in 44.8% (94/210) of CSF samples of the patients and culture in 8.1% (17/210) while 85.7% (180/210) yielded mDNA concentrations of ≥ 0.005ng/ml. Of the 180 samples, spectrophotometric reading for DNA purity value of ≥ 1.7–12.20 (good quality DNA) was recorded in 55 (30.6%), 1.0–1.69 (quality DNA) in 103 (57.2%), 0.57–0.99 (low quality DNA) in 14 (7.8%) with only 8 samples (4.4%) failing purity evaluation (with value of 0.00 at A260/280). In a recent metagenomic study by Zhang et al., (32) on 135 patients, 26 (19.3%) were culture positive while 32 (23.7%) were identified by metagenomic next generation sequencing (mNGS). This is the closest method to the one we used. While their method was mNGS, ours is still at the level of mDNA extraction from CSF samples and subsequent molecular identification and characterization by qPCR of the three bacteria of interest (N. meningitidis, H. influenzae, and S. pneumoniae). Our study therefore provides a strong baseline data for processing CSF samples for qPCR without the need for culture, and shows that majority of samples would yield quality DNA material (> 1.0 ng/µL).

The gel electrophoresis results showed spatial bands, that could be linked to the percentage of agarose in the gel used (1.5%), which is good in resolving linear DNA molecules size range of 300–3000 bp (31) as against the amplicon size of the bacteria of interest; N. meningitidis (127 bp), H. influenzae (113 bp), and S. pneumoniae (51 bp). The strength of our research lies on the fact that metagenomic protocol does not rely on bacterial culture and isolation for extraction of mDNA for use in the detection of aetiological agents of meningitis, but rather on the constituent DNA concentration present in the sample. However, one limitation to our study is that we did not include viral pathogens, being that RNA was not extracted. Another limitation is that we did not perform restriction digest of the extracted mDNA or PCR amplification of 16S rDNA on the extract, which would have confirmed the suitability of the extracted mDNA for downstream processing.

Conclusion:

Quality mDNA from CSF samples will ensure successful qPCR results for rapid and accurate detection of bacterial pathogens in meningitis, beginning first at molecular detection using multiplex real-time PCR (rt-PCR) down to species-specific singleplex rt-PCR. This will eliminate the time and labour consuming traditional culture methods often associated with “yielded no growth syndrome” or abysmal low yield of CSF culture output resulting in the very poor outcome of laboratory confirmed cases of cerebrospinal meningitis (CSM).

Acknowledgements:

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Authors contributions:

PIC and IEI conceived, and led the design and writing of the manuscript. PIC, NUP, UYB, DLD, OCN, and MR processed the CSF samples at the various sites for phenotypic methods, storage, and transportation of the samples to Abuja and finally to Safety Molecular Pathology Laboratory, Enugu for storage at ~80°C. NE and NCR were responsible for molecular biology techniques (PCR) orientation, training and processing of CSF samples extraction and quality check of DNA. PIC fully participated in performance of the PCR activities. PIC, IEI, and NE were responsible for the final editing of the manuscript.

References:


