Molecular Characterization of Measles Virus among Children in Parts of North Western Nigeria

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A B S T R A C T

Molecular characterization of measles virus among children in parts of north western Nigeria, out of 30 samples analysed for PCR with specific primers, 21 samples confirmed presence of measles virus infection. Genotype B3 strain was confirmed in four (4) samples selected from different location within the state in Northern Western Nigeria. Sokoto, Jigwa and Kaduna States, Ten (10) IgM positive samples from each States were randomly selected and subjected to RT PCR, out of the ten (10) IgM positive samples from each state. Twenty (66.7%: 20/30) IgM positive samples shows a band with an amplicon size of 527bp. Four measles virus were subjected to sequencing and BLAST and all were genotype B3 strain. The BLAST information obtained from the queried sequences and the closest corresponding sequences from the GenBank corresponded with genotype B3 from Genbank, The phylogenetic relatedness of measles virus in this study with those in the GenBank revealed a common homology.

Keywords: Measles virus infection, Aerosolized secretions, Systemic infection, Nasopharynx, viremia.

Introduction

Measles virus belonging to the genus Morbillivirus in the family PARAMYXOVIRIDAE is an enveloped virus with a non-segmented, negative-strand RNA genome (Griffin, 2001). It is 100 – 200 nm in diameter and two membrane envelope proteins are important in pathogenesis which are the F (fusion) protein, which is responsible for fusion of virus and host cell membranes, viral penetration, and hemolysis, and the H (hemagglutinin) protein, which is responsible for adsorption of virus to cells (Bellini et al., 2005). The World Health Organization (WHO) currently recognizes 8 clades designated A, B, C, D, E, F, G, and H. Within these clades, there are 23 recognized genotypes, designated A, B1, B2, B3, C1, C2, D1, D2, D3, D4, D5, D6, D7, D8, D9, D10, E, F, G1, G2, G3, H1, and H2, and one provisional genotype, d11 (WHO, 2015).

In order to infect a cell, a virus must first bind to a cellular receptor on the surface and enter the cell. The presence of such a receptor determines whether the cell is susceptible to the virus. Measles is a highly infectious disease characterized by fever, respiratory symptoms, redness of the eyes and a
complications are common and may be quite serious. The rash usually starts on the head and then spreads to the rest of the body. Fever can persist, reaching extremely high temperatures, rash can last for up to a week, and coughing can last about 10 days (Jawestz et al., 2011).

The causative agent of measles virus is generally transmitted by aerosolized secretions deposited on upper respiratory tract mucosal surfaces. Exposure leads to local respiratory tract replication; infection of regional lymphoid tissues then occurs followed by viremia and systemic dissemination as revealed by the characteristic skin rash. Most children recover uneventfully from the illness, but serious complications can occur, including pneumonia and involvement of the central nervous system (Parks et al., 2009). Airflow studies demonstrated that droplet nuclei generated in the examining room used by the source patient were dispersed throughout the entire office suite. Airborne spread of measles from a vigorously coughing child was the most likely mode of transmission (Bloch et al., 2005).

Measles is a systemic infection and the primary site of infection is the respiratory epithelium of the nasopharynx. Two to three days after invasion and replication in the respiratory epithelium and regional lymphnodes, a primary viremia occurs with subsequent infection of the reticuloendothelial system. Following further viral replication in regional and distal reticuloendothelial sites, a second viremia occurs 5–7 days after initial infection. During this viremia, there may be infection of the respiratory tract and other organs. Measles virus is shed from the nasopharynx beginning with the prodrome until 3–4 days after rash onset (Gerber et al., 2009). Prevention of measles infection rests on successful immunization with currently available live, attenuated vaccine. The immunity produced by the vaccine lasts many years and is probably life-long. The recommended age of administration in infant immunization programmes is 9 to 15 months. Vaccine efficacy is 85% at 9 months of age and increases to 90–95% at 12–15 months of age. Measles virus is highly transmissible. High routine immunization coverage can reduce measles incidence but will not prevent accumulation of susceptible individuals, which can lead to outbreaks if virus is introduced into a population where the number of susceptible individual are above the critical threshold for that population (WHO, 2010).

The use of vitamin A in treatment has been investigated. A systematic review of trials into its use found no significant reduction in overall mortality, but it did reduce mortality in children aged less than two years (Souza et al., 2002).

Five out of six WHO regions have set goals to eliminate measles, and at the 63rd World Health Assembly in May 2010, delegates agreed to a global target of a 95% reduction in measles mortality by 2015 from the level seen in 2000, as well as to move towards eventual eradication. However, no specific global target date for eradication has yet been agreed (WHO, 2010).

Materials and Methods

Study Area

The study was conducted in some North Western region of Nigeria, Sokoto, Jigawa and Kaduna State.

Study Population

Children 0-12 years of age with suspected measles symptoms like, fever of 38°C or greater, cough, redness of the eyes and
maculopapular rash, attending selected hospitals in the study area were included.

Sample Size

A convenience sampling technique was employed; therefore, 30 blood samples of children attending some selected hospital in the North western Nigeria were collected.

Sample Collection and Analysis

Blood samples were collected aseptically from each clinical case child by an experienced technician and dispensed into sterile labeled plain specimen bottles.

RNA Extraction

Total measles virus RNA from clinical samples was obtained using the QIAamp Viral RNA kit (Qiagen Hilden Germany) according to manufacturer’s protocol. Blood samples from measles cases were obtained by centrifugation for 2mins at 2300rpm. RNA was extracted from pellet using the spin-column procedure. Two hundred microlitre (200ml) of each positive blood sample was transferred into a clean tube. Five hundred microlitre (500µl) of lysis buffer AVL was dispensed into each clean tube and incubated at room temperature for 10 minutes. The samples were mixed by pulse – vortexing for one minute. The sample mixtures were centrifuged at 12000rpm for five minutes. The spin columns were transferred into a clean 2µl collection tube and the filtration was discarded. Five hundred microlitre (500ml) of wash buffer AWI was added to each spin column and centrifuge at 12000rpm for two minutes. The filtrate was discarded. The empty spin columns were centrifuged at 10000rpm for two minutes. The spin columns were transferred into new clean tubes and 60µl of elution buffer AVE was added into each spin column and allowed to stand at room temperature (25°C) for 2 minutes. Centrifugation was done at 10000 rpm for two minutes to elute the RNA from the spin columns into the tube. A double elution was done by adding 100ml of elution buffer AVE into each spin column and centrifuged after two minutes at 10000rpm for two minutes. RNA samples were stored at – 80°C until required for use.

RT-PCR Amplification

Specific cDNA of Measles Virus nucleoprotein was synthesized by a reverse transcription using Superscript III Reverse Transcriptase (Invitrogen, Merelbeke, Belgium) and random hexamers (Invitrogen). The Measles Virus cDNA was amplified by nested PCR using primers (Table 1) MN5 ( 5′ - GCC ATG GGA GTA GGA GTG GAA C - 3′) and MN6 ( 5′ - CTG GCG GCT GTG TGG ACC TG - 3′ ) for the first round and primers Nf1a ( 5′ - CGG GCA AGA GAT GGT AAG GAG GTC AG -3′) and Nr7a (5′- AGG GTA GGC GGA TGT TGT TCT GG -3′) for the second round. Both PCRs were performed in a total volume of 25 μL that contained 1.8 mmol/L MgCl$_2$, 1× PCR buffer, 0.2 mmol/L dNTPs, 0.5 U Platinum Taq (Invitrogen), and 0.8 µmol/L forward and reverse primer (Eurogentec, Seraing, Belgium). One microliter of cDNA or 5 µL of first-round product (diluted 50x in water) was added as template. Cycling conditions was denatured at 94°C for 2 min; 35 (first round) or 30 (second round) cycles of amplification at 94°C for 30sec, 55°C (first round) or 58°C (second round) for 1 min, and 72°C for 1 min; and a final extension at 72°C for 5 min

Detection and Identification of the Amplified PCR Products

Agarose gel of 1% concentration was prepared by weighing one gram of agarose in a conical flask and adding 100ml of 1x TAE
buffer (Tris Acetate EDTA). The agarose was solubilized by heating in a microwave oven for two minutes. The solubilized agarose was allowed to cool to 40\(^\circ\)C and 5ml of ethidium bromide was added. The gel was poured into the gel tray with the combs in place was left to solidify for about 50 minutes. Electrophoresis tank was filled with the running buffer (Tris Acetate EDTA) and the gel slab was totally submerged in the buffer at a level not more than two millimeters above the gel. The combs were gently removed to give well-formed wells. About 20ml of each amplified PCR product was loaded into respective wells and 100 volts was applied for 30 minutes using Bio–Rad power pack (powerpac TM HC). The power supply was turned off after 30 minutes. The gel was removed and examined using Bio–Rad Gel Doc Molecular Imager System (Gel Doc TM XRT). The basepair of the amplicon was determined using the standard (100bp DNA ladder).

**Cycle Sequencing**

Sequencing reaction were Prepare in a 0.2mL thin-wall tube with 2 tubes per template DNA for a forward sequencing primer and a reverse sequencing primer reaction. All reagents were kept on ice box while preparing the sequencing reactions, 8 μl of sterile water and 2 μl of diluted plasmid DNA (at 100 ng/ul) were added to each sequencing tube. The tubes were spanned briefly. The plasmid water was preheat and mixed at 96\(^\circ\)C for 1 min in the thermo cycler and then cools on ice box. 8 μl DTCS Quick Start Master was added and Mixed to each tube followed by 2 μl of T-7 forward sequencing primer added to one of the tubes for a template and 2 μl of SP6 reverse sequencing primer was added to the other tube for a template.

The reaction components were mixed thoroughly. The liquid at the bottom of the tube was consolidated and briefly centrifuging before thermo cycler with the following conditions; 96\(^\circ\)C for 20 seconds, 50\(^\circ\)C for 20 seconds and 60\(^\circ\)C for 4 minutes

**Cycle Sequencing Clean up**

Nested PCR products were purified by using the Jetquick PCR product Purification Spin Kit (Genomed, Lohne, Germany). Twenty-five cycles of cycle sequencing (2-min elongation) were performed by using a BigDye Terminator version 6.1 Cycle Sequencing kit (Applied Biosystems, Nieuwerkerk, the Netherlands) with Nf1a or Nt7a primers (0.5 μmol/L) and 10 ng of purified PCR product. The product was sequenced for measles virus genotyping using an ABI 3130 Genetic Analyzer (Applied Biosystems). Phylogenetic trees were constructed by using the neighbor-joining method (Kimura 2-parameter) and MEGA7 version software

**Results and Discussion**

Ten (10) samples from each States were subjected to RT-PCR. 10 samples selected from Sokoto State, 6 samples selected from Jigawa State and 4 samples selected from Kaduna State showed bands for measles virus. (Plate I, II, and III). Four (4) measles virus isolate (products of RT-PCR) from the studied states were sequenced, 2, 1 and 1 isolate from Sokoto, Jigawa and Kaduna State respectively. All the 4 sequenced were genotype B3 strain. The distribution of measles virus in the respective states in the study area is presented in tables 1, 2, 3 and 4.

The BLAST information obtained from the Query sequences and the closest corresponding sequences from the genbank revealed that samples S3, S4, J21 and K24 correspond with genotype B3 sequences from the genbank. This is presented in table 5 and 6.
**Table 1** Detection of measles virus from blood samples in Sokoto state

| S/N | Sample Number | RT-PCR |
|-----|---------------|--------|
| 1   | S3            | +      |
| 2   | S4            | +      |
| 3   | S19           | +      |
| 4   | S24           | +      |
| 5   | S25           | +      |
| 6   | S26           | +      |
| 7   | S50           | +      |
| 8   | S51           | +      |
| 9   | S70           | +      |
| 10  | S78           | +      |

Key: S = sample number from Sokoto state, + = positive product of RT PCR, - = negative product of RT PCR

**Table 2** Detection of measles from blood samples in Jigawa State

| S/N | Sample Number | RT-PCR |
|-----|---------------|--------|
| 1   | J5            | -      |
| 2   | J10           | +      |
| 3   | J21           | +      |
| 4   | J29           | -      |
| 5   | J66           | -      |
| 6   | J76           | +      |
| 7   | J82           | +      |
| 8   | J92           | +      |
| 9   | J98           | +      |
| 10  | J102          | -      |

Key: J = Samples in Jigawa State

**Table 3** Detection of measles virus from blood samples in Kaduna state

| S/N | Sample Number | RT-PCR |
|-----|---------------|--------|
| 1   | K2            | -      |
| 2   | K3            | -      |
| 3   | K8            | +      |
| 4   | K24           | +      |
| 5   | K28           | -      |
| 6   | K36           | -      |
| 7   | K40           | -      |
| 8   | K43           | +      |
| 9   | K63           | -      |
| 10  | K76           | +      |

Key: K = Samples Collected from Kaduna
Table 4 Percentages prevalence of measles virus in the north western Nigeria

| State     | n  | % occurrence |
|-----------|----|--------------|
| Sokoto    | 10 | 10(10)       |
| Jigawa    | 10 | 6(60)        |
| Kaduna    | 10 | 4(40)        |
| **Total** | **30** | **20(66.7)** |

Key: n= Number of IgM Positive Sample; Number in Parentheses= Percentage % = Percentage

Table 5 Cycle sequencing products

| Location | MV Genotype B3 Sequencing                                                                 |
|----------|------------------------------------------------------------------------------------------|
| SOK3     | TTAGGGGTGTCCGTGTCTGAGTCTTGTCTTTAGATTTGCATGTCCTGACCATGGCTTGAGCCTGAACTCCGATGCA          |
|          | GAGCTGGCTGGAGTCCCCGACTGTAAGTGGTCTTGACGCTCTGCTGGCTGAGTCCTGCACCTGACTGGAATGCTCACCTCTTC |
| SOK4     | CATTTGACACTCTAGGGGTGTCCGTGTCTGAGTCTTGTCTTTCCGAGATTGCATCTGGCA                      |
|          | TGGCTGGAGCTACGGAGGGGACTGCTTGAGCTTGGGGACTGCTGGCCTGGCTGAGTGGCTGCAGGTGCTGACTGGAATTC |
| JG21     | TTAGGGGTGTCCGTGTCTGAGTCTTGTCTTTAGATTTGCATGTCCTGACCATGGCTTGAGCCTGACCTGAGCACCTG    |
|          | GAGCTGGCTGGAGTCCCCGACTGTAAGTGGTCTTGACGCTCTGCTGGCTGAGTCCTGCACCTGACTGGAATGCTCACCTTC |
| KD24     | CGAGTTAAGCCGAGCTTCCGAGATGCAAGTGGTCTTGAGTCTTGTCTTTAGATTTGCATGTCCTGACCATGGCTTGAGC |

KEY: Sok3= Sokoto sample number 3, Sokoto4 = Sokoto sample number 4, JG21 = Jigawa sample number 21, KD24 = Kaduna sample number 24
Plate 1 Agarose Gel Electrophoresis of RT-PCR products of Measles virus IgM Positive Samples from Sokoto States

Key: Lane M 100bp DNA Ladder; Lane 1-10 amplicons products; Lane + = Positive Control; Lane - = Negative Control
Plate.2 Agarose Gel Electrophoresis of RT-PCR products of Measles virus IgM Positive Samples from Jigawa and Kaduna States

Key: Lane M 100bp DNA Ladder; Lane 11, 14, 15, 20, 21, 22 and 25 were negative; Lane 12, 13, 16, 17, 18, 19, 23 and 24 shows amplicons of Measles virus; Lane - = Negative Control

Table.6 Closest Affiliates of Measles Virus Sequence Obtained Compared with those from GenBank

| Seq | Accession No  | Max Score | Total Score | Query length | E – value | Identity |
|-----|--------------|-----------|-------------|--------------|-----------|----------|
| S3  | KX838946.2   | 764       | 764         | 442          | 0.0       | 98%      |
| S4  | KX949610.1   | 722       | 722         | 443          | 0.0       | 98%      |
| J21 | KX949603.1   | 773       | 773         | 436          | 0.0       | 99%      |
| K24 | KT732217.1   | 785       | 785         | 450          | 0.0       | 99%      |

The query sequences covers almost 100%

Plate.3 Agarose Gel Electrophoresis of RT-PCR products of Measles virus IgM Positive Samples from Kaduna State

Key: Lane M 100bp DNA Ladder; Lane 26, 27 and 29 were negative; Lane 28 and 30 shows amplicons of Measles virus; Lane + = Positive Control; Lane - = Negative Control
Phylogenetic tree were prepared using Mega 7 version software and the neighbor – joining method with 1000 bootstrap replicates. Sequences from North Western Nigeria Measles virus strains were compared with the sequences of B3 genotype on Genbank. This is presented in figure 3.

Twenty (66.7%) of the 30 IgM positive samples were positive for measles virus gene. This agree with the finding of Mohammoud et al., (2013), Zhang et al., (2012), who reported the incidences of measles virus in Africa and China respectively. This could be due to long period of sampling before analysis because the virus may not be able to survive a long period of time before screening, but the IgM antibody will still be present in the sample no matter the period it stayed before screening. This result also shows that RT – PCR is a sensitive and specific assay for detecting measles virus RNA in a variety of clinical samples. This agrees with the finding of Jangu et al., (1995) who detected measles virus RNA in clinical samples using RT-PCR. The entire four measles virus were subjected to sequencing and BLAST. All the four were genotype B3 strain, this agrees with the finding of Michaela et al., (2005) who reported that Genotype B3 was first detected in 1993 in Gambia but have subsequently been detected in cases from Cameroon, and Central Africa and has been imported into numerous countries including France, Germany and the USA. Genotype B3 is clearly the endemic genotype in most parts of the African Continent where it is widely distributed (WHO, 2012).

The BLAST information obtained from the queried sequences and the closest corresponding sequences from the GenBank revealed that samples S3 and S4 from Sokoto State corresponded with genotype B3 from Genbank, Sample J21 and K24 from Jigawa and Kaduna State respectively also corresponded with B3 strain in the GenBank.

The phylogenetic relatedness of measles virus in this study with those in the GenBank revealed a common homology. Measles virus S3 and S4 on the same leave node were genetically highly related to one another and related to those gotten from Calais (FRA/01:16) and Sokoto (NGA/20.13), but vary with those on a neighbouring clade positions. Measles virus J21 is genetically closely related to Measles virus S4 on the neighboring leave node while Measles virus K24 is genetically closely related to Measles virus J21 on the neighboring clade position. This could be due to evolutionary relatedness, this could be that, they all emolliated from the same ancestor.

In conclusion, the measles virus strain detected in this study by sequencing and phylogenetic analysis was genotype B3 Strain and the phylogenetic analyses in this study revealed a close relationship of the query sequences with the one obtained from France, Newcastle, London and North central part of Nigeria (Niger).

Acknowledgement

We wish to express our deep sense of gratitude to the management of Sokoto, Jigawa, and Kaduna State Ministry of health for giving us the ethical approval to carry out this research. We also appreciate the support rendered by staff at various facilities.

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How to cite this article:
Abdul Fatai, K., O.S. Olonitola, M. Aminu and Jatau, E.D. 2017. Molecular Characterization of Measles Virus among Children in Parts of North Western Nigeria. Int.J.Curr.Microbiol.App.Sci. 6(7): 3981-3991. doi: https://doi.org/10.20546/ijcmas.2017.607.412