INTRODUCTION

The types of influenza virus found in pigs are known as swine influenza generally called swine flu or swine-origin influenza virus (S-OIV) (1, 2). Influenza virus belongs to the genus Orthomyxovirus in the family Orthomyxoviridae which consists of influenza A, B and C viruses and has an envelope, single-stranded, negatively sensed RNA, eight separate segments and pleomorphic appearance with an average diameter of 120nm (3, 4).

Influenza A virus causes swine influenza which is an acute, highly contagious respiratory disease (5) with classical aetiological types that include influenza A subtypes H1N1, H1N2, H2N3, H3N1 and H3N2 and rarely influenza C while influenza B has not been reported in swine (6, 7, 4). This is because type A group continually undergoes antigenic shift and drift unlike B and C which are relatively stable (8, 4). Generally, 16 haemagglutinin (H or HA) and 9 neuraminidase (N or NA) subtypes have been identified. This means pigs can also be infected with other subtypes as they play a substantially important role in the ecology of influenza A virus and can act as a “mixing vessel”. When co-infections among human, avian or swine influenza viruses occur within a specific host, a new subtype can be produced by antigenic re-assortment (9, 10).

The virus usually spreads via aerosols produced by an infected person when coughing, sneezing or spitting or through contaminated hand to eye, nose or mouth either from fomites or direct personal contact such as hand-shake(11, 12).

In Nigeria, influenza viruses have caused a significant amount of morbidity in the general population but the incidence of excess mortality is unknown. Until the recent advancement in the field of molecular virology, the detection of influenza virus has always been by isolation method (13). This study therefore evaluates the detection of influenza A virus in order to establish that different strains can co-circulate among local pigs in Lagos which can cause possible intermittent infection in man.

MATERIALS AND METHODS

Study site

The study subjects were collected from pigs at two different sites in Ayedoto farm settlement, Agric, Ojo Local Government, Lagos between June and September, 2010.

Sample population, collection and preservation

A total of 58 (58 nasal and 58 throat=116 samples) apparently healthy land race pigs aged between 2-30 months old were the subject for this study. Nasal and throat swab samples that contained adequate numbers of ciliated and columnar epithelial cells were collected from land race pigs. Each swab was transferred into a total of 58 (58 nasal and 58 throat=116 samples) apparently healthy land race pigs aged between 2-30 months old were the subject for this study. Nasal and throat swab samples that contained adequate numbers of ciliated and columnar epithelial cells were collected from land race pigs. Each swab was transferred into commercially available sterile cryovials containing 2ml aliquot of Dulbecco’s modified eagle medium transport medium with antibiotic to prevent desiccation, death and bacterial growth. They were conveyed to the laboratory in coolers with dry ice packs immediately after collection. The samples were kept at 4°C for extraction the next day (< 24hrs) since freezing and thawing reduce the ability to recover virus. Aliquots of samples were frozen at -70°C.

Extraction Process

ABSTRACT

This study detected and subtyped strains of influenza virus from pigs in Lagos, South-western Nigeria. A total of 116 (58 nasal and 58 throat) samples from healthy pigs were analysed from two different sites in Ayedoto farm at Ojo Local Government between June and September, 2010 using reverse transcription polymerase chain reaction (RT-PCR). Influenza virus type A 31(26.7%) was detected. Subtyping was done using RT-PCR with H1, H3 and H5 primers and only subtypes H1 [5(16.1%)] and H5 [5(16.1%)] were detected. No positive detection was made for subtype H3. This research work is the first documented detection of influenza A virus in pigs in Lagos, Nigeria and demonstrates the need for a sustainable surveillance mechanism of swine and other influenza viruses to be able to prevent influenza epidemic in the environment.

Keywords: Subtype, Influenza A, Pig, Lagos

DETECTION OF INFLUENZA A VIRUS IN PIGS IN LAGOS, NIGERIA

RUNNING TITLE: INFLUENZA A VIRUS IN NIGERIA

Anjorin, A. A. *, Omilabu, S. A. 2, Salu, O. B. 3, Oke, B. O. 4

1Department of Microbiology, Lagos State University, Ojo, Lagos, Nigeria 2Department of Medical Microbiology and Parasitology, College of Medicine, University of Lagos, Ile-Ife, Osun, Nigeria 3Department of Medical Microbiology and Parasitology, College of Medicine, University of Lagos, Ile-Ife, Osun, Nigeria 4Department of Human virology, Microbiology Division, Nigerian Institute of Medical Research, Yaba, Lagos

*Correspondence: Anjorin, A. A. Email: azlaboratories@yahoo.co.uk

http://dx.doi.org/10.4314/ajcem/v13i1.4

AFR. J. CLIN. EXPER. MICROBIOL 13(1): 41-45...
This study extracted RNA genetic material by diatomaceous sand method using Qiagen kit (Germany). 140ul of sample was added to 560ul of lysis buffer with carrier RNA by dispensing each into sterile Eppendorf vial followed by addition of 100 mg diatomaceous sand (Sigma-Aldrich, USA) in order to trap and bind the RNA (5). 560ul of absolute ethanol was added and mixed by pulse vortexing until homogeneous mixture was obtained. Incubation was done at room temperature for 20 minutes with vortexing at 5 minutes interval. The mixture was then spun at 13,000 rpm for 1 minute while the supernatant was decanted. Serial washings using two kinds of washing buffer (AW1 and AW2: 500ul each) were carried out before each centrifugation at 13,000 rpm for 1 minute and separation of supernatant. 400 ul of acetone was later added with another vortexing and centrifugation at full speed 13,000 rpm for 2 minutes. The supernatant was then separated and stored in RNA 1.5ul Eppendorf tube at -80°C before amplification.

**Master Mix Preparation and PCR Amplification**

A one step process was used for both reverse transcription and PCR amplification treatment process with Qiagen (Germany) one step kit. The master mix used for one reaction before the addition of 5ul of suspected influenza RNA extract for the synthesis of complementary DNA (cDNA) include: 5ul Rnase free water, 5ul 5x RT buffer, 5ul Q-solution, 1 uldNTP-mix, 1.5 ul each of forward and reverse primer and 1 ul enzyme mix. The RT-PCR mixture was then incubated at 50°C for 30 minutes for reverse transcription followed by 45 cycles run in a thermo cycler (Eppendorf, Germany) PCR machine. Denaturation - 95°C for 15 minutes; Activation - 95°C for 30 seconds; Annealing - 55°C for 30 seconds; Elongation - 72°C for 30 seconds; and Extension- 30°C for 30 seconds (14, 15, 5).

**Agarose Gel Electrophoresis Process**

Identification of amplicons was carried out by agarose gel electrophoresis method with 2% agarose gel (Peolab, Germany). 2 ul of SYBR gel stain (Invitrogen, USA) was added. The gel was later poured into a casting block with inserted comb. It was allowed to solidify for 15 minutes. The comb was gently removed while the gel was placed inside an electrophoresis tank containing TAE buffer 50x (Genaxxon, Germany). 1ul of 6x loading dye (Fermentas, EU) plus 5ul of amplicons were loaded per lane onto the gel. 3ul of a 100bp DNA ladder (Invitrogen, USA) was loaded along with the samples as marker. Electrophoresis was carried out in a Westburg electrophoresis machine (Biometra, Netherland) for 30 minutes by 3-4cm at 130 volts. Ultra violet (UV) source safe imager trans illuminator (Invitrogen, USA) was used to illuminate and viewed the gel pictures before they were taken by Biodoc analyze 2.0(Biometra, Germany). The correct RNA amplification was indicated by correct size and a comparison with positive and negative controls of influenza A (figure 1) and subtypes H1 and H5 (figure 2).

**RESULTS**

A total of 116 (58 nasal, 58 throat) samples from 58 pigs were studied. Influenza A virus 31(26.7%) was detected using RT-PCR (Table 1). The 31(26.7%) positive influenza A virus detected were typed with influenza A subtypes H1 [5(16.1%)] and H5 [5(16.1%)] results obtained. No positive detection was made for subtype H3 (Table 2).

| TABLE 1: DISTRIBUTION OF INFLUENZA A VIRUS USING RT-PCR METHOD |
|------------------|------------------|------------------|
| INFLUENZA TYPE   | NO OF SAMPLES TESTED | NO OF POSITIVES (%) |
| A                | 116              | 31(26.7%)        |

| TABLE 2: DISTRIBUTION OF SUBTYPES OF INFLUENZA A VIRUS USING H1, H3 AND H5 PRIMERS |
|-----------------------------------|------------------|------------------|
| INFLUENZA A SUBTYPE               | NO OF SAMPLES TESTED | NO OF POSITIVES (%) |
| H1                                | 31               | 5(16.1%)         |
| H3                                | 31               | -                |
| H5                                | 31               | 5(16.1%)         |
Figure 1: Amplicons of Influenza A virus separated by Agarose Gel Electrophoresis (2%) for 30 minutes by 3-4cm at 130 volts: Panel A has seven total no of positives as shown on lane 1, 4, 11, 12, 13, 17 and 18 when compared with the positive control (P) on last lane as ruled by the straight line passing through the center; Panel B has five total no of positives (57, 65, 68, 70 and 72); Panel C. No of positives = 6 (76, 78, 81, 85 and 86); Panel D. No of positives = 13 (100, 101, 102, 104, 105, 106, 107, 108, 109, 111, 113, 114 and 116).
Figure 2: Amplicons of Haemagglutinin (H) 1 and Haemagglutinin (H) 5 protein genes of subtypes of Influenza A virus separated by Agarose Gel Electrophoresis (2%) for 30 minutes by 3-4cm at 130 volts. Panel A: Subtype H1; No of positives = 4 (1, 2, 4 and 5); Panel B: Continuation of Subtype H1; No of positives = 1 (53); Panel C: Subtype H5; No of positives = 5 (49, 50, 51, 52 and 53). Negative (N) and Positive (P) controls are as shown on second to the last and last lane respectively. 100 base pair Ladder (L) was used as the marker as shown on each gel panel.

DISCUSSION

Influenza virus is notoriously known for its unique ability to cause recurrent influenza epidemics and global pandemics during which acute febrile respiratory illness occurs explosively. There are two qualities of influenza virus that account for much of its spread. First is the ability to emerge and circulate in avian or porcine reservoirs by either genetic reassortment or direct transmission and subsequently spread to human at regular intervals. Second, is the fast and unpredictable antigenic change of important immune target once the virus is established in human (16).

The high number of positive results to influenza A virus 31(26.7%) detected in this study is characterized by its continuous antigenic drift and shift, making it more genetically diverse with high prevalence in the subject (8,4). This is further supported by the work of (17) which showed that influenza A virus mutates at a rate 2-3 times faster than type B which is relatively stable.

Only sub types H1 [5(16.1%)] and H5 [5(16.1%)] detected in this study disagree with the work of (1) and (5) that detected subtype H3 in pigs. (18) and (19) also detected and worked on both H1 and H3 Thailand and European subtypes in pigs respectively. No positive detection of H3 in this study may be due to H3 primer mismatch to local strains of influenza A in Nigeria since foreign primer (Qiagen, Germany) were used which produced unclassified bands.

The detection of H5 agrees with the work of several authors since H5 subtypes are commonly found in birds (20). Another reason is that birds are being reared side by side with the pigs at the site of this study. Subtyping in this study however did not include the use of neuraminidase (N) primers due to cost and in availability in Nigeria.

The ability of RT-PCR to detect influenza virus types agree with the work of (21) that RT-PCR is capable of detecting the virus even when the viral genomes are present in low level and is generally more sensitive in the detection of influenza virus than any other method including serology and culture (22).
REFERENCES


