Seasonal emergence of swine erysipelas in hilly state Nagaland, Northeast India

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Seasonal outbreaks of swine erysipelas have been reported in back yard pig farms in the Phek district of Nagaland, India. The alpha haemolytic isolate of *Erysipelothrix rhusiopathiae* was recovered on blood agar from the clinical samples. The organisms were confirmed microscopically, biochemical analysis as well as by polymerase chain reaction (PCR) amplification of 16S rRNA gene and sequence analysis. These Nagaland isolates (KT160358, KT160359) were closely related to the type sp. *E. rhusiopathiae* in phylogenetic analysis and forms the same clad with Chinese isolates of swine and murine origin indicating an epidemiological link. The isolates were found to be most sensitive to oxytetracycline and responded to treatment. Swine erysipelas occurred in Phek district in a season due to sudden change of weather and temperature. Pigs exposed to such predisposing factors probably favoured to propagation of already persisted organisms in pigs. This is the first confirmed case of *E. rhusiopathiae* infection from the NE states of Nagaland, India.

Key words: Swine erysipelas, *Erysipelothrix rhusiopathiae*, pig, polymerase chain reaction (PCR) Nagaland, India, Oxytetracycline.

INTRODUCTION

*Erysipelothrix rhusiopathiae*, belonging to the family *Erysipelotrichaceae*, is a non-motile, Gram-positive, non-sporulating, non-acid-fast organism distributed worldwide affecting wide variety of vertebrate and invertebrate species including man (Reboli and Farrar, 1989). Organisms in many occasions harbour by pigs in lymph nodes and shed along with feces, urine, saliva and nasal secretions (Lee et al., 2011). Affected pigs manifest the disease as (i) acute septic form, (ii) subacute urticarial form marked by reddish-purple rhomboid spots or "diamonds" in the skin, (iii) joint or arthritic form, and (iv) chronic cardiac form (endocarditis) (Reboli and Farrar, 1989). Various predisposing factors, change of environmental conditions and parasitic infestation lead to...
Outbreaks of Swine Erysipelas – samples collection

The disease of swine erysipelas was reported from Porba village (altitude 1856 MSL), District Phek of Nagaland during the summer rainy season from 2013 to 2015. The village had a pig population of 552 cross bred and 200 local doom pigs as per the livestock census report 2012 (GOL, 2012). The pigs were kept in the small yard small unit mainly for meat purpose and fed them on household as well as hotel waste. In every rainy summer season (May – July) there was disease outbreaks in pigs. Affected animals (150) clinically were anorexic and recumbent with high fever (106° F) for 2-3 days. Some animals developed erythematous patches. Postmortem examination revealed haemorrhages in intestine, congestion of liver, spleen and kidney. During the course of investigation 3 more animals died within 12 days. All affected pigs were also reported to be infested with pig louse Haematopinus suis. Clinical sample like swabs from the wound and tissue biopsy from the affected areas were collected in sterile containers for isolation of the causative organism. Samples also included sloughed off tissues and biopsy samples preserved in 10% formalin for histopathological study.

Identification and antibiotic activity of Erysipelothrix rhusiopathiae

The swab samples (N=51) from the affected areas were inoculated into nutrient broth and incubated aerobically at 37°C for 48 h. Sub-culturing was done on blood agar plates, incubated at 37°C in presence of 5% CO2 for 24 h. Colony morphologies were studied. Gram’s staining and biochemical analysis were done to confirm the organism. A panel of antibiotic discs containing amikacin (30 µg), amoxicillin (30 µg), ampicillin (10 µg), cefotaxime (30 µg), chloramphenicol (30 µg), ciprofloxacin (5 µg), cloxacillin (30 µg), cotrimoxazole (25 µg), enrofloxacin (5 µg), gentamicin (10 µg), neomycin (30 µg), norfloxacin (10 µg), oxytetracycline (30 µg), streptomycin (25 µg) and tetracycline (30 µg) was used to study the antibiotic sensitivity pattern of the isolates. The zone of inhibition was measured, recorded and interpreted according to the clinical and Laboratory Standards Institute criteria (CLSI-MIC).

Detection of nucleic acid of Erysipelothrix rhusiopathiae and sequencing

For detection of E. rhusiopathiae nucleic acid tissue samples were processed using DNA Sure Tissue Mini Kit (Nucleo-pore, cat.#NP-61305) and polymerase chain reaction (PCR) amplification for 16S rRNA gene primers (MO101 F AAGATGCCATAGAAACGCTTGA and M0102 R CTGTAATTGCCCATATACTA) of E. rhusiopathiae (Makino et al., 1994) was used. The PCR conditions were optimized with a final volume of 25 µl at 94°C for 5 min followed by 30 cycle of 94°C for 30 s, 54°C for 2 min and 72°C for 45 s and final extension was carried out at 72°C for 5 min. In PCR reaction 28 ng/µl total genomic DNA was taken along with positive and negative control. Products of PCR were visualized in 2% agarose gel electrophoresis under Gelsoc (Kodak, USA). Further PCR products were purified by QIAquick PCR purification kit protocol and sequenced.

Analysis of gene sequence and phylogenetic studies on Erysipelothrix rhusiopathiae

To determine the relationship of the Nagaland isolates of E. rhusiopathiae with other isolates of this species, 16S rRNA gene was amplified and sequenced (GenBank accession numbers KT160358 and KT160359). Sequence identity at nucleotide level was determined by Clustal W method of Meg-Align program in DNA Star package (DNASTAR Inc., USA). The phylogenetic tree was constructed using other 16S rRNA gene sequences available at NCBI (viz. strain ZYL(KF811052.1), isolate EU188793.1, strain KG-B1(AB055309.1), strain Fujisawa(NP_074878.1), isolate DO462571.1, strain: ATCC 19414(AB055905.1), strain JPBP25120905(HM569359.1), strain Chiba9393(EEF49748.1). sp. TI27.5(JG739693.1), sp. LV19(KJ670316.1), strain Er.GXL-1(KP063151.1), inopinata strain 143-02 (inopinata strain 143-02) strain: JCM 8534(LC019778.1) and strain Er.GXBY-1(KP063149.1).

Histopathology

Formalin fixed tissues were processed as per standard protocol for histopathological studies. Sections of 4-5 micron thickness were stained routinely with Haematoxylin and Eosin stain and observed under oil immersion objective of a low power light microscope.

RESULTS AND DISCUSSION

Tissue samples processed for detection of classical swine fever virus and swine pox virus were confirmed as negative for both viral agents. However, bacteriological investigation demonstrated association of E. rhusiopathiae infection in affected (5) as well as in dead (2) pigs. Prevalence of erysipelothrix in many animals, mostly in pigs and birds has been reported throughout the world including from India (Shankar et al., 2009; Arora et al., 2011). But there was a single report on swine erysipelothrix (SE) from Meghalaya (Das et al., 2014). North Eastern Region has the highest pig population of the country. Diverse geographical locations, varied climatic situations and frequent movement of pigs favour for spread of the disease through carrier pigs (Leslie et al., 2015). Present report is a thorough investigation on swine erysipelothrix occurred seasonally at Nagaland, another NE state of India.

Clinical signs

A total of 150 pigs during 2013-15 exhibited clinically high fever (105-107°F), anorexia, firm faeces, and animals...
lying down. Cutaneous lesions appeared after 5-7 days of illness initiating with erythematous patches followed by papules of 4-5 mm diameter, dark purple, raised, firm to touch giving square to rhomboid shape in entire body. No vesicular stage was noticed. Although scabs developed in the entire body, prominent lesions were identified on head, face back, belly, limbs, tail and on ear (Figure 1). In unattended cases scab lesions were sloughed off within 17-22 days leaving a large ulcerated area. These typical cutaneous lesions have been mostly seen in grower animals. In affected pigs it appears that cutaneous lesions were developed following acute stages of illness.

Considering the pathogenesis of *E. rhusiopathiae*, organisms gain access to the body, probably through the tonsils or other lymphoid tissue of the digestive tract and spread throughout the body. The bacteria produce neuraminidase, an enzyme that cleaves mucopolysaccharides in cell walls which may mediate the widespread vascular damage that accompanies SE. Vascular damage leads to thrombosis and interference with microcirculation in capillaries and venules at many sites. Classic cutaneous rhomboid urticaria (diamond skin) occurs in a percentage of pigs shortly after the acute febrile stages. In younger pigs with acute erysipelas, signs are similar, with cyanosis of extremities, ears and snouts pronounced and urticaria less common (Anonymous, 2016).

Seasonal occurrence of swine erysipelas in Nagaland justifies the stress due to sudden change of temperature. As stated by Amanda (2012) that stress factors such as overstocking, mixing pigs after weaning, and sudden changes in temperature can trigger clinical erysipelas.

### Isolation and identification of organism

On blood agar the organism produced small, circular and transparent α-hemolytic colonies with a smooth glistening surface and edge. Biochemically all the isolates were catalase, oxidase and urease negative, produces H₂S and ferments glucose and lactose. All 7 isolates were identified as *E. rhusiopathiae* based on the cultural, morphological and biochemical characteristics. The Nagaland isolates were sensitive in vitro to oxytetracycline, tetracycline, ampicillin, amoxicillin and cloxacillin; moderately sensitive to streptomycin, enrofloxacin, amikacin, co-trimoxazole, cefotaxime and ciprofloxacin; and resistant to gentamicin and norfloxacin, chloramphenicol and neomycin. Based on the antibiogram, survived ailing pigs were treated with oxytetracycline at 10 mg/kg body weight, intramuscularly for one week. Eight out of 13 treated animals responded promptly and recovered. Treatment of swine erysipelas cases with penicillin (Shankar et al., 2009) or other penicillin group of drugs such as the combination of amoxycillin and cloxacillin (Das et al., 2014) have been frequently reported.

### Molecular confirmation and characterization

Tissue samples were negative for Swine fever virus and pox virus in PCR. A total of seven *E. rhusiopathiae* isolates recovered from tissue samples were subjected for PCR amplification using 16s rRNA gene specific primer set. All isolates were found to be positive for *E. rhusiopathiae* with amplification products of 407bp.
Figure 2. 2% agarose gel showing PCR Products. Here, L1-100 bp marker, L2 NTC, L3 & 4 PCR products.

Figure 3. Phylogenetic tree of Nagaland isolates of *E. rhusiopathiae* (KT160358, KT160359 as compared to the ATCC 19414 type strain and other isolates.

Two Nagaland isolates KT160358 and KT160359 were sequenced and showed high sequence identity with the sequences of other *E. rhusiopathiae* available in the GenBank database. The Nagaland isolates (accession numbers KT160358, KT160359) were in the same clad (Figure 3) along with other strains originated from China. Phylogeny based on the nucleotide and amino acid sequences of the viruses provides a better understanding of the molecular epidemiology of the isolates. The state
Nagaland shares international boundaries with Myanmar and having territorial link with other neighboring international countries like China, Bhutan and Bangladesh. The Chinese strains used in the construction of phylogenetic tree viz. swine isolates KPO 63150.1, strain Er.GXBY-1(KP063149.1), strain Er.GXL-1(KP063151.1) and murine isolate DQ462571 are forming same clad with Nagaland isolates (KT160358 and KT160359) indicated an epidemiological link. Movements of animals and animal products might spread the infection to this locality.

Histopathology

Histopathologic alteration of affected skin showed extensive damage to the capillaries and venules of dermis with infiltration of lymphoid cells and fibroblasts. Congestion and presence of microthrombi in the capillaries were consistent finding in all skin biopsy samples investigated in the present study (Figure 4). Similar observations were also recorded by Shankar et al. (2009).

Cultural characteristics, molecular confirmation and histopathological changes conclusively proved that swine erysipelas is prevailing in Nagaland. Pigs harbouring the infection manifest clinically at sudden change of climate. It is estimated that 30–50% of healthy swine harbour the organism in their tonsils and other lymphoid tissues (Stephenson and Berman, 1978). Again, trans border movement of pigs and their products could facilitate spreading of SE in this locality. Further study on prevalence of swine erysipelas and identification of carrier pigs can provide actual guidelines to control the disease in this part of India.

Conclusion

Unorganized pig farms in the Phek district of Nagaland experienced high mortality of grower pigs during rainy-summer season. A detail isolation and molecular investigation confirmed the association of swine erysipelas in this part of north eastern region of India. This is the only report on seasonal occurrence of swine erysipelas in the hilly low temperate climate of the NE states of Nagaland, India.

Conflict of Interests

The authors have not declared any conflict of interests.

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