Full Length Research Paper

Isolation and identification of *Archanobacterium pyogenes* (*Actinomyces pyogenes*) from Arabian gazelles

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*Archanobacterium pyogene* (*Actinomyces pyogene*) is an opportunistic pathogen of economically important livestock such as dairy, beef cattle and gazelles. It is also a common inhabitant of the mucous membranes of these animals. This study was aimed to investigate the epidemiology of *A. pyogenes* in the infected Arabian gazelles kept at King Khalid Wildlife Research Center at Thumamah, 70 km north of Riyadh, Saudi Arabia and the relationship between symptoms and infection by *A. pyogenes*. Samples were collected from pleural fluid of 70 infected gazelles and from upper respiratory tract of 30 healthy animals. Enrichment and isolation from the collected samples resulted in 48 bacterial strains, 8 strains from upper respiratory tract of healthy animals and 40 from infected gazelles. Biochemical tests of the 48 isolated bacterial strains revealed that all the strains were beta-hemolytic on blood agar, hydrolyzed gelatin and starch were unable to hydrolyze esculin or reduce nitrate and did not produce urease which was typical characteristics of *A. pyogene*. Identification of the isolated strains was further confirmed by molecular method using analysis of 16S rDNA.

Key words: Opportunistic, pathogen, wildlife, Arabian gazelles, Actinomyces pyogenes.

INTRODUCTION

*Archanobacterium pyogene* (*Actinomyces pyogene*) is an anaerobic Gram-positive bacilli and it is a part of the normal flora in many domestic animals (Carlos Emilos et al., 2009). *A. pyogenes* is a rare cause of pyogenic infections in humans and in most of cases; reports are questionable since there was failure to definitively identify the pathogen (Hermida Ameijeras A et al., 2004). *Gazelles* belong to Kingdom Animalia, Phylum, Chordata, Class, Mammalia, Order, Artiodactyla, Family, Bovidae, and Genus Gazelle. Three species of gazelles are considered native to the Kingdom of Saudi Arabia including sand gazelles *Gazella subgutturosa marica*, Arabian mountain gazelles, *Gazella gazelle* and the Saudi gazelle *Gazella dorcas* (saudiya). However, the distribution of gazelles in Saudi Arabia has declined dramatically during recent decades due to excessive hunting and habitat degradation. Today, Gazelles survive only in small numbers in a few isolated parts of their former range. Therefore, King Khalid Wildlife Research Center (KKWRC) was established in 1986 by the National Commission for Wildlife Conservation and Development (NCWCD) to breed native Arabian *Gazelles* species as part of the conservation program for the Kingdom of Saudi Arabia. The center was developed at Thumamah, 70 km north of Riyadh, on the farm of the late King Khalid, whose animal collection formed the basis of the breeding stock. One year later, the management of the animal collection was given to the Zoological Society of London on behalf of the (NCWCD). Different species of gazelles kept at the center are for breeding, research and reintroduction purposes. Two main species of gazelles are kept at the KKWRC including, *G. gazella*, locally known as Idmi and *G. subgutturosa marica*, locally known as Reem. Previous study identified, determined the susceptibility and detected gene cassettes of *A. pyogenes* isolates from cows with endometritis (Liu, et al., 2009). It is a commensal and an opportunistic pathogen of
economically important livestock, causing diseases as diverse as mastitis, liver abscessation and pneumoni (Ceribasi, et al., 2009). Clinical and pathological features of three clinical cases of testicular abnormalities associated with A. pyogenes, cellulites and testicular degeneration, confirm that A. pyogenes can be a testicular pathogen under field conditions (Gouletsou et al., 2006). Intrauterine inoculation of nulliparous ewes with A. pyogenes and Escherichia coli produced an antibody response and reduced the severity of subsequent infections (Meghan et al., 2005). A. pyogene possesses a number of virulence factors that contribute to its pathogenic potential and expresses a cholesterol-dependent cytolysin, pyolysin, which is a haemolysin and is cytolytic for immune cells, including macrophages (Jost and Billington, 2005). Therefore, more insight on pathogenic sequence of A. pyogenes is curial for understanding its physio-pathogenicity. The sequence data from autosequencers, mining Web-databases, performing automatic and manual sequence alignment, analyzing sequence facilitates the promotion and better understanding of the underlying assumptions (Tamura et al., 2007). Accurate identification of bacterial isolates is an essential task in clinical microbiology. Fatal A. pyogenes endocarditis case in human, mostly related to living in rural areas and to contacts with animals has been described by DNA sequence (Plamondon et al., 2007). However, phenotypic methods are time-consuming and either fails to identify some bacteria such as Gram-positive rods entirely or at least fail to do so in some clinical situations (Bosshard et al., 2003). The traditional identification of bacteria on the basis of phenotypic characteristics is generally not as accurate as identification based on genotypic methods (Claridge et al., 2004).

Recombinant DNA (rDNA) sequencing is an effective means for the identification of aerobic Gram-positive rods which are difficult to identify by conventional techniques. 16S rDNA sequencing is a recent method of identification which offers a useful alternative (Mignard and Flandrois, 2006). Partial 16S rDNA was amplified (by using primers p27f and BAC1401r and sequenced by using primers 1100r, 765fs and 10f. Sequences were compared with those available in GenBank using gapped BLASTN 2.0.5 software (Carlos et al., 2009). The objectives of this study was to investigate the epidemiology of A. pyogenes in the infected Arabian gazelles under captive at the KKWRC, investigation of the relationship between symptoms and infection by A. pyogenes, and determination of the virulence factors encoding genes of the isolated A. pyogenes. It is usually recommended to compare this molecular identification tool to classical phenotypic methods. The great potential of the method has been reported for Gram-positive rods and coryneform bacterial identification (Bosshard et al., 2003). Therefore, 16S rDNA gene sequencing was used in this study to confirm the identification of the bacterial strains isolated from different infected and healthy gazelles.

**MATERIALS AND METHODS**

Materials used include captive healthy gazelles at King Khalid Wildlife Research Center, breeding pens, infected gazelles, pleural fluid gazelles, syringes and transport media, as well as specimens of upper respiratory tract of 30 healthy gazelles, 15 Idmi and 15 Reem gazelles.

**Reagents and experimental instruments**

Sterile saline, 5% sheep blood agar medium and Brain heart infusion (BHI) agar media, sterile loop, 5% CO2, Trypticase Soy Broth medium (TSB), 5% fetal bovine serum, 10% (v/v) sterile glycerol, API Coryne kit (Biomerieux, France), soft agar media, conventional Gram stain, distilled water, buffered glutaraldehyde solution (2.5% glutaraldehyde in 0.05 M sodium cacodylate buffer pH 7.4), phosphate buffer, osmium tetroxide solution (1.0% osmium tetroxide in 0.05 M sodium cacodylate buffer, pH 7.4) and graded ethanol series (25, 50, 75, and two 100% each for 10 min; 5 × 10 = 50 min). In addition, swabs coated with gold, methanolic uranyl acetate solution, Reynolds lead citrate, mineral oil, GP medium, Blood & Tissue Kits (Qiagen, NY, USA), 0.2 μM nucleopore filters. For DNA sequencing, the following were used: proteinase K, ethanol 96-100%, 20 mM Tris-Cl (pH 8), 0.2 mM sodium EDTA, 1.2% Triton® X-100, lysozyme, agarose, loading dye solution (Qiagen, NY, USA), One kb DNA ladder (Roche, Mannheim, Germany), ethidium bromide solution (1 μg/ml), 50x TAE (Tris-Acetate/EDTA) buffer stocking solution, universal eu bacterial forward primer 16F27 (5'-AGA GTT TGA TCC TGG CTG AG-3') and reverse primer 16R152S (5'-AAG GAG GTG ATC CAG CCG CA-3'), MWG Biotech AG (Ebersberg, Germany), deoxynucleoside triphosphates, MgCl2, Taq polymerase and 10x PCR buffer. More also, instruments used include: dissecting microscope, scanning electron microscope (SEM), Transmission Electron Microscope (TEM), pH indicator, DNeasy Mini spin column, micro centrifuge tube, vortextes, pipette.

**Isolation of A. pyogene**

Specimens were collected from pleural fluid of 70 infected gazelles. Specimens were also collected from upper respiratory tract of 30 healthy gazelles (15 Idmi and 15 Reem gazelles). Samples were kept in cold and transferred directly to the lab within 1 h for further analysis. Enrichment and isolation of A. pyogenes was carried out according to the methods reported by (Ertas et al., 2003). Pleural fluid samples from the abscessed lungs were serially diluted in sterile saline solution up to 10^-4. Aliquots (200 μL) and swabs (collected from infected and healthy gazelles) were inoculated (in triplicates) onto 5% sheep blood agar medium and Brain heart infusion (BHI) agar media supplemented with 5% sheep blood using sterile loop and incubated aerobically and in 5% CO2 at 37°C for 72 to 96 h. The obtained single colonies were sub-cultured several times in fresh agar plates until pure homogeneous single colonies were obtained. The cells of the isolated bacteria were then grown in the trypticase Soy Broth medium (TSB) supplemented with 5% fetal bovine serum and incubated at 37°C for 18 h.

Identification of the isolated bacteria was carried out using conventional methods, including morphological, biochemical tests, API Coryne kit (Biomerieux, France) and was finally confirmed by molecular methods using analysis of 16S rDNA sequence as previously reported. Colony morphology of 24- to 48-h old cultures was examined under a dissecting microscope. Hemolytic activity was recorded on 5% sheep blood agar media after incubation at 37°C for 72 h. Cell motility was examined using soft agar media according to the method described by (Harmon et al., 2005). 3 μL of
overnight culture of the isolates were inoculated carefully to avoid any disturbance of the agar surface, on the center of soft agar medium (0.4%). The plates were then incubated at 37°C and monitored up to 72 h. Motile bacteria migrated from the plate center and diffuse into the medium causing turbidity, whilst non-motile bacteria show growth confined to the center.

Identification of cell morphology

Cell morphology and cell wall type were examined using conventional Gram stain and the scanning electron microscope. The cell morphology of the bacterial isolates was examined under SEM in the Electron Microscopy Unit available in the National Research Center, Faculty of Science, King Saud University as follows (Allan-Wojtas et al., 2010): The API Coryne kits contained API Coryne strips, GP medium, McFarland standard, nitrate A, nitrate B, Zym A, Zym B, PYZ, H2O2, mineral oil and sterile saline 3 ml. The procedures of strip inoculation and results interpretation were carried out according to the manufacturer’s instructions and the results were interpreted according to the API guideline.

Gene sequencing analysis

The isolated bacterial strains were further identified using 16S-rDNA sequence analysis. First, the total bacterial DNA was extracted, then the 16S-rDNA gene was PCR amplified, sequenced and the obtained sequences were compared with the sequences of known bacteria in the GenBank database.

DNA extraction and purification

All the isolates were inoculated into TSB medium supplemented with 5% fetal bovine serum and incubated overnight at 37°C. 3 ml of growth cultures were centrifuged and the pellets were washed twice with sterile saline solution. The total bacterial total DNA was extracted using DNeasy Blood & Tissue Kits (Qiagen, NY, USA) with a resultant yield of 10 μg DNA. The total extracted DNA was subjected to electrophoresis with 1 × TAE in agarose gel by loading (12 μL) mixed with 3 μL of loading dye solution (Qiagen, NY, USA) into the gel wells. One kb DNA ladder (Roche, Mannheim, Germany) was used as the molecular size standard. The gel was placed in ethidium bromide solution (1 μg/ml) for 30 min and finally was placed on an ultra violet transilluminator to visualize the DNA (Ausubel et al., 2008). To extract the PCR product from the agarose gel, the gel was washed twice with sterile distilled water and placed on an ultra violet transilluminator to visualize the DNA and the amplified 16S-rDNA products were sliced off from the agarose with sterile razor blade.

The DNA was purified from the agarose using QIAquick gel Extraction Kits. The 16S-rDNA gene of the isolated strains (n = 48) were sent for sequencing to Eurofins MWG Operon Sequencing Facility (Fraunhoferstr, Martinsried, Germany) using the same forward and reverse primers described earlier (Volker et al., 1996). The obtained sequences of 16S-rDNA of the isolates were aligned with the reference 16S-rDNA sequences of the European Microbiological Laboratory (EMBL), GenBank (gb, Germany) and the Database of Japan (DDBJ) that are available in the National Centre for Biotechnology information (NCBI).

RESULTS

Figure 1 shows health gazelles at the KKWRC, breeding pens. A necrotic cavity containing an air-fluid level was recoreded. The infection extended into the pleural space and produced an empyema without rupture of the abscess cavity. The necrotizing pneumonia was restricted to one pulmonary segment or lobe, and in few cases it encompassed an entire lung or both lungs (Figure 2). Microscopic examination of the infected lungs tissues revealed similar supplicative lesions in all of the infected gazelles. Pulmonary changes consisted of multifocal-to-coalescing areas of necrosis, associated with cell debris and fibrin, bounded by low numbers of neutrophils. A direct smear from necrotic areas showed Gram-positive bacilli forming chains (Figure 3). Enrichment and isolation of A. pyogenes from the collected samples resulted in the isolation of 48 bacterial strains. The isolates showed colony surrounded with narrow sharp hemolytic zone, morphology similar to those reported for A. pyogenes (Figure 4).

Furthermore, cell morphology and size were examined under (SEM) (Figure 5) and Transmission Electron Microscope (TEM) (Figure 6). Cell length and diameter were about 0.8 to 1.0 μm and 0.3 μm, respectively. The bacterial isolates were subjected to further identification by different biochemical tests. API strip identified all isolated bacterial strains (n = 48) as A. pyogenes. The results also indicated that all isolates were non-motile, beta-hemolytic on blood agar, able to hydrolyze gelatin and starch, and in addition, the isolates were unable to

PCR amplification of 16S-rDNA gene

16S rDNA was amplified by PCR using universal eubacterial forward primer 16F27 (5’-AGA TTG TGA TCC TGG CTC AG-3’) and reverse primer 16R1525 (5’- AGG GTG ATC CAG CCG CA-3’) derived from E. coli 16S-rDNA sequence synthesized by MWG Biotech AG (Ebersberg, Germany). The PCR amplification was performed using the purified genomic DNA as a template. The total volume of the PCR reaction was 50 μL (Forward primer, 1 μ (10 pmol/μL), reverse primer, 1 μ (10 pmol/μL), deoxynucleoside triphosphates 0.6 μL (10 mmol/L), 1 X PCR buffer 3 μL (10 x), MgCl2 21.8 μL (25 mM), Taq polymerase 0.2 μL (5 U/μL), DNA template 2.5 μL (100 ng) and sterile aqua dest water (19.9 μL). The PCR reaction was performed in a Peltier thermal cycler (BioRad, USA) using the following conditions: Initial denaturation 95°C for 5 min, (Denaturation 95°C for 30 second, Annealing 52°C for 30 second, Extension 70°C for 1.5 min, final extension 70°C for 5 min) 34 cycles) and End/store Forever 4.

Agarose gel electrophoresis of the PCR products

The PCR products (12 μL) were mixed with 3 μL of loading dye solution (Qiagen, NY, USA) and were separated in a 1% (w/v) agarose gel electrophoresis at 120 V in 1 μL × TAE buffer with a 1 kb DNA ladder (Roche, Mannheim, Germany) as the molecular size standard. The gel was placed in ethidium bromide solution (1 μg/ml) for 30 min and was placed on an ultra violet transilluminator to visualize the DNA (Ausbubel et al., 2006). To extract the PCR product from the agarose gel, the gel was washed twice with sterile distilled water and placed on an ultra violet transilluminator to visualize the DNA and the amplified 16S-rDNA products were sliced off from the agarose with sterile razor blade.

The DNA was purified from the agarose using QIAquick gel Extraction Kits. The 16S-rDNA gene of the isolated strains (n = 48) were sent for sequencing to Eurofins MWG Operon Sequencing Facility (Fraunhoferstr, Martinsried, Germany) using the same forward and reverse primers described earlier (Volker et al., 1996). The obtained sequences of 16S-rDNA of the isolates were aligned with the reference 16S-rDNA sequences of the European Microbiological Laboratory (EMBL), GenBank (gb, Germany) and the Database of Japan (DDBJ) that are available in the National Centre for Biotechnology information (NCBI).
Figure 1. Postmortem examination of infected lungs of the gazelle.

Figure 2. Results of the Gram staining of the isolates; isolated strains were gram positive short bacilli.

Figure 3. Isolates grown on 5% sheep blood agar medium at 37 °C for 24-72 h Strains; showed colony surrounded with narrow sharp hemolytic zone, morphology similar to those reported for Actinomyces pyogenes.
hydrolyze esculin or reduce nitrate and did not produce urease. However, there were considerable variation in other biochemical tests particularly sugars fermentation. The results of the biochemical tests of 11 isolates out of 48 are presented in Table 1.

Genomic DNA of the isolated was extracted using DNeasy extraction kit as described earlier. The concentration and purity of the purified bacterial DNA was determined by measuring OD at 260 and 280 nm. Good total DNA yield in the range of 11.7 µg to 27.5 µg was obtained. The DNA purification was repeated for any sample with OD 260/280 ratio higher than 2.0. The extracted DNA was further checked using agarose gel electrophoresis (Figure 7). The amplified 16S rDNA was...
Table 1. The results of the biochemical tests of 11 isolates out of 48 isolates.

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Fermentations of:

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Xylose                       |          | +    | +    | -    | +    | +    | -    | +    | +    | +    | -     | +     |
Mannitol                     |          | -    | -    | -    | -    | -    | -    | -    | -    | -    | -     | -     |
Maltose                      |          | -    | +    | +    | -    | -    | +    | -    | -    | +    | -     | +     |
Lactose                      |          | +    | +    | +    | +    | +    | +    | +    | +    | +    | +     | +     |

Figure 6. Extracted DNA from *Actinomyces pyogenes* isolates under agarose gel electrophoresis.

The 16S rDNA of the isolated strains was amplified by conventional PCR using universal eubacterial forward primer (16F27) and reverse primer (16R1525), and the PCR products were analyzed using agarose gel electrophoresis and visualized using ultra violet (Figure 9). The sequence of 16S rDNA (length...
Figure 7. Amplified by 16S rDNA of the isolates by conventional PCR using universal eubacterial forward primer (16F27) and reverse primer (16R1525) under agarose gel electrophoresis.

Figure 8. The amplified 16S rDNA sliced off from the agarose gel under electrophoresis.

= 1020 bp) of strain KGD5 has been deposited in the GenBank database with accession number of GU358474 and designed as *A. pyogenes* KGD5 and could be reached using the following website: http://www.ncbi.nlm.nih.gov/nuccore/283553395 (Figures 10 to 19).

**DISCUSSION**

Many of suppurative lung infections of the captive gazelles have been recorded at KKWRC. Actinomyces pyogenes is a common inhabitant of the mucous membranes of the upper respiratory, gastrointestinal, and
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**Figure 9.** Alignment of 16S rDNA sequence of KGH1 isolate. Identities = 818/832 (98%) with *Arcanobacterium pyogenes* isolate HC-H 10 16S ribosomal RNA gene, strand=Plus/Plus. Yellow color: KGH1 sequence.
Figure 10. Alignment of 16S rDNA sequence of KGH2 isolate. Identities = 940/946 (99%) with Arcanobacterium pyogenes isolate BM-H 07-1 16S ribosomal RNA gene, strand=Plus/Plus. Yellow color: KGH2 sequence.
Figure 11. Alignment of 16S rDNA sequence of KGH3 isolate. Identities = 869/875 (99%) with *Arcanobacterium pyogenes* isolate HC-H 10 16S ribosomal RNA gene, strand=Plus/Plus. Yellow color: KGH3 sequence.
Figure 12. Alignment of 16S rDNA sequence of KGD4 isolate. Identities = 763/771 (98%) with Arcanobacterium pyogenes isolate BM-H 07-1 16S ribosomal RNA gene, strand=Plus/Plus. Yellow color: of KGD4 sequence.
Figure 13. Alignment of 16S rDNA sequence of KGD5 isolate. Identities = 1015/1020 (99%) with *Arcanobacterium pyogenes* HC-H 10 16S ribosomal RNA gene, strand=Plus/Plus. Yellow color: KGD5 sequence.
Figure 14. Alignment of 16S rDNA sequence of KGD6 isolate. Identities = 621/627 (99%) Arcanobacterium pyogenes isolate HC-H 10 16S ribosomal RNA gene, strand=Plus/Plus. Yellow color: KGD6 sequence.
urogenital tracts of a number of domestic animal species (Jost et al, 2005). The objectives of this study were to investigate the epidemiology of *A. pyogenes* in the infected gazelles under captive at King Khalid Wildlife Research Center, investigation of the relationship between symptoms and infection by *A. Pyogenes*. Cell morphology and size were examined under SEM and TEM. Cells were about 0.8-1.0 µm and 0.3 µm in length.

**Figure 16.** Alignment of 16S rDNA sequence of KGD8 isolate. Identities = 601/603 (99%) with *Arcanobacterium pyogenes* isolate HC-H 16S ribosomal RNA gene, strand=Plus/Plus. **Yellow color:** KGD8 sequence.
Figure 17. Alignment of 16S rDNA sequence of KGD9 isolate, Identities = 774/783 (98%) with Arcanobacterium pyogenes isolate HC-H 10 16S ribosomal RNA gene, strand=Plus/Plus. Yellow color: KGD9 sequence.

and diameter respectively. The isolated bacterial strains (n = 48) were subjected to identification by biochemical testes and API Coryne strip. The biochemical tests revealed that all isolates were beta-hemolytic on blood agar, able to hydrolyze gelatin and starch, in addition, the isolates were unable to hydrolyze esculin or reduce
nitrate and did not produce urease which are typical characteristics of *A. pyogenes* according to Bergey's Manual of Systematic Bacteriology (Farmer III, et al., 2005). However, there was some variation in sugar fermentation ability of different isolates which are also similar to the results reported for *A. pyogenes* by other scientists who reported that different strains of *A. pyogenes* showed variable fermentation ability of sugars (Jost et al, 2005).

The results of the bacterial isolation of samples collected from pleural fluid of infected gazelles indicated that *A. pyogenes* was detected in 57% of the infected gazelles (40 out of 48 pleural fluid samples). The identification of the isolated strains was further confirmed by molecular method using analysis of 16S rDNA gene sequencing. The PCR products were analyzed using...
agarose gel electrophoresis and purified from the agarose gel. Purified 16S-rDNA fragment was sequenced twice and were obtained up to 1020 bases with minimal mistakes. The 16S-rDNA sequences of the isolated bacterial strains were compared and aligned with the known 16S-rDNA sequences of other bacteria available in the GenBank database. It was found that the isolated bacterial strains had 98% to 100% similarity with different strains of \textit{A. pyogenes}. It has been reported that 16S-rDNA gene sequences similarity of \( \geq 97\% \) indicated the same bacterial species (Muyzer et al., 1993). Therefore, the isolated strains, to best of our knowledge are the first

\textbf{Figure 19.} Alignment of 16S rDNA sequence of KGD11 isolate. \textit{Arcanobacterium pyogenes} isolate HC-H 11 16S ribosomal RNA gene.
report of \textit{A. pyogenes} isolation from infected Arabian gazelles in Saudi Arabia.

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