Clinical and microbiological aspect of surgical affections associated with current epidemics in bovine.

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Abstract

Egypt is endemic with Lumpy Skin Disease (LSD) and Foot and Mouth Disease (FMD), both impose a drastic effect on the economy. These viral diseases are often complicated with secondary bacterial infections; however, scarce data is available. In this study, utilizing microbiological and molecular procedures, we identified a number of secondary bacterial infections complicating FMD- and LSD-infected cases. Moreover, the antimicrobial resistance profile of the bacterial isolates was explored. The data showed that several bacterial pathogens including Serratia odorifera, Enterococcus faecium, Paenibacillus apiaries and Microbacterium hominis could be recovered from LSD-infected cases. On the other hand, FMD cases were found to be complicated with Pseudomonas mosselii, Bacillus cereus and Trueperella pyogens. However, Bacillus licheniformis was isolated from both disease conditions. The antimicrobial testing revealed that all pathogens are resistant to more than three antibiotics of different families, indicating the multi-drug resistant capabilities of these isolates. This baseline investigation confirms the presence of many secondary bacterial infections linked to LSD and FMD as well as the widespread antimicrobial resistance among the isolated pathogens, however, the contribution of these pathogens to the mortality rate in both conditions needs further studies.

Keywords: lumpy skin disease, foot-and mouth-disease, antimicrobial, 16S rRNA sequencing.
INTRODUCTION

Epidemic attacks in cattle have been reported to be widely spread in various disease forms all over the world. In Egypt, the most endemic diseases challenging livestock and causing severe financial losses on the national scale are foot-and-mouth disease (FMD) and lumpy skin disease (LSD) (Bauer. 1997 and Murply et al. 1999). Several deleterious effects, such as lowering the productivity and reproductively of affected animals and death were recorded (Fubini and Ducharme 2016). In addition, the costs and efforts consumed in supportive care and treatment of such complications, especially the surgical one, are alarming (Kossaibati and Esslemont 1997).

LSD was first reported in Egypt in May 3, 1988 (Fayed, 1988 & Salem, 1989). It occurred in Suez province and the source of infection was imported cattle from Somalia. The causative agent of LSD is a Neethling virus, genus Capri pox virus in the family poxviridae (Murpaly et al, 1999). There is only one report of the natural occurrence of LSD in species other than cattle, water buffalo (Bubalis), but no further cases are recorded (House, 1998 and Radostits et al, 1994).

An outbreak of LSD infection in different ages and sexes of Egyptian cattle with severe and serious complications was recorded in the summer of 2006, in many Egyptian governorates. Clinical signs, epidemiological characters, electron microscopy and histopathological findings of LSD in this outbreak were reported by Fayez and Ahmed(2011). The skin of different parts of infected animals body such as muzzle, nares, back, limbs, scrotum, perineum, eyelids, ears, nasal and oral mucosa and the tail displayed nodules 1-7 cm in diameter. The nodules were painful and involved the epidermis, dermis and subcutaneous tissue and further involve the musculature. With the progresses of the disease, the nodules became necrotic with a deep scab (sit fast) forms (El –Kanawaty, 1989; Ismail 1989 and House 1998). Lesions on the udder and teats resulted in mastitis, teat fistula or obstruction (Heidrich and Renk 1967; Ismail 1989 and House 1998). Lameness also may result from secondary bacterial infection. Also, severe edema in the brisket, ventral abdomen and limbs was evident (El-Kanawaty, 1989; Kenawy, 1991 and House 1998).

Foot-and-mouth disease virus (FMDV) is a member of the genus Aphthovirus. There are seven major viral serotypes of the virus: O, A, C, SAT 1, SAT 2, SAT 3 and Asia. The most common serotype worldwide is serotype O. It is responsible for an Asian pandemic that affected many countries throughout the world began in 1990 (Bauer, 1997). However, serotype C is uncommon and was not reported since 2004 (Loth et al., 2011). While other serotypes caused serious outbreaks, some FMDV serotypes are more variable than others. However, they may collectively contain more than 60 strains with the potential of development of new strains (Bauer, 1997 and Salt, 1998). While most of the strains affect all susceptible host species, some have a more restricted host range (e.g., serotype O Cathay strain, which only affects pigs). Immune response of animals infected with one FMDV serotype does not protect it from other serotypes infection. Defense against other strains within the same serotype depends on their antigenic determinant similarity (Salt, 1998). Vesicular lesions are the most characteristic feature of FMD (Bauer, 1997).

Since the 1950s, several outbreaks of the FMD affected cattle, buffaloes, sheep,
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goats and camels, with the prevalence of serotype O1 FMD virus (FMDV). Therefore, attention has been drawn to the economic importance of FMD in Egypt (Moussa et al. 1974, Daoud et al. 1988 and El-Nakashly et al. 1996). In 2006, the last outbreak due to serotype (O) was reported by Abd El-Rahman et al. (2006). Since 1972 serotypes other than (O) have not been reported (Aidaros, 2002). Clinical cases of FMD were recognized on a cattle farm in Ismailia, Northeastern Egypt in 2006. Laboratory examination was carried out for clinical samples to determine FMD serotype by using enzyme linked immunosorbent assay (ELISA) test, and reverse transcriptase polymerase chain reaction (RT-PCR). FMDV type (A) was detected and designated as A/Egy/2006 (Knowles et al. 2007). Since 2006, the bivalent (FMD) vaccine, containing both O1 and A/Egy/2006, was locally produced. After this incident in July 2006, reported FMD outbreaks are caused by either serotype (O1 and A) (GOVS, 2009 and Ghoneim et al., 2010). Mastitis or hoof malformations with accompanying signs of pain are most common complications of FMD infection. In addition, sudden death of young calves due to heart failure without developing vesicles was also reported (Ahmed et al. (2012) and Kandeil et al (2013).

Bobkurov (1977) and Frank (1981) stated that knowledge of bacteriology should be used in diagnosis and treatment of surgical infected lesions. They mentioned that the pyogenic bacteria are the most important in acute surgical infections. Micro-organisms are the real causes of epidemic diseases and their complication, either (virus, bacteria, fungi). There are infectious and non-infectious micro-organisms, so isolation and identification of micro-organism must be done in proper conditions to ensure the validity of the testing procedure, avoid false positive results due to contamination, and to prevent the spread of infectious microbial agents (EL- Mishad, A (1991)). Therefore the aims of the work are to study the clinical picture of the complications of some current epidemic diseases and management. We attempt to isolate and identify the causative agents of these complications and to perform antibiotic sensitivity tests to choose the best antibiotic to be used to treat bacterial diseases.

MATERIALS AND METHODS

The present study was carried out on the prevalent clinical and microbiological aspect of surgical affections in bovine in Qena governorate. Most of the cases were registered at the village hospitals and at the owners’ houses in the period between April 2018 and April 2020 on a total number of 60 animals (bovine). All cattle were raised on small holding groups and were cross breeds. The study included both sexes and ages from 2 to 4 years, table (1). Clinical findings were systemically identified and evaluated for its nature and severity. Sterile swabs were used to collect samples from different lesions from random 10 cases of LSD (five cases) and Foot FMD (five cases).

Microbiological Study:

1- Collection of samples:

Sterile cotton swabs were used to collect samples from different sites (infected nose, mouth, wound, eye, udder and claw) and transferred immediately on ice to our microbiology lab (Department of Microbiology, Faculty of Veterinary Medicine) for microbiological study.
2- Isolation and identification of microorganisms:

Sterile cotton swabs were submerged in tryptone soy broth (lab M) and incubated at 37 °C for 18-24 hour to enhance the growth of microorganism. After incubation, a loopful of each bacterial suspension were streaked on different agar plate (including; blood agar (trypton soya agar + 5% sheep blood), MacConkey agar (Oxiod) and mannitol salt agar (Himedia) and incubated at 37 °C for 24-48 hours with and without 5% CO₂. The macroscopic morphology of the colonies, hemolysis and pigment production were identified according to Quinn et al., 1994 and Sears et al., 1993. Identification of bacteria was carried out by gram staining to detect the microscopical morphology of the bacteria (Forbes et al 2002 and Bebora et al 2007) then subjected to biochemical identification (oxidase test, catalase test and coagulase test). The suspected bacterial isolates were subjected to 16S rRNA sequencing to identify genus and species.

3- PCR Identification to bacterial species.

3.1. DNA extractions from bacterial isolates were done according to QIAamp DNA mini kit manufacture instructions.

3.2. PCR amplification. The amplified 16S rRNA gene was obtained from each isolate and reference strain by PCR with the universal primers F27 (5-AGAG TTTGATCMTGCTCAG-3) and R1492 (5-TACGGYTACCTTGTTAC GACTT-3) (Lagacé et al., 2004), which are targeted to universally conserved regions and permit the amplification of an
approximately 1485 bp fragments. PCR amplification was carried out in Thermal cycler (Biometra). Reaction tubes contained 25 μL (6μL template DNA, 12.5 μL of Emerald Amp GT PCR master mix (Takara), 1 μL of each forward and reverse primer (20 pmol) and 4.5μL of PCR grade water). Initial DNA denaturation and enzyme activation steps were performed at 94 °C for 15 min, followed by 35 cycles of denaturation at 94 °C for 30s, annealing at 56 °C for 1 min and extension at 72 °C for 1 min, and a final extension at 72 °C for 10 min. The presence and yield of specific PCR product (16S rRNA gene) was monitored on 1% agarose (BioShop) gel electrophoresis at 200 V for 1 hour in 1x Tris-acetate-EDTA buffer and made visible by ethidium bromide staining and UV transillumination according to Sambrook et al., 1989.

4- Sequencing of 16S rRNA.

4.1 Purification of the PCR Products were done according to manufacture instruction of QIAquick gel extraction kit (Qiagen).

4.2 Sequencing reaction:
A purified PCR product was sequenced in the forward and reverse directions on an Applied Biosystems 3130 automated DNA Sequencer (ABI, 3130, USA). Using a ready reaction Bigdye Terminator V3.1 cycle sequencing kit. (Perkin-Elmer/Applied Biosystems, Foster City, CA). A BLAST® analysis (Basic Local Alignment Search Tool) (Altschul et al., 1990) was initially performed to establish sequence identity to GenBank accessions. The sequence reaction was done according to the instruction of the manufacture as follows in table (2).

4.3. Phylogenetic analysis:
A comparative analysis of sequences was performed using the CLUSTAL W multiple sequence alignment program, version 1.83 of MegAlign module of Lasergene DNAStar software. Pairwise, designed by Thompson et al., (1994), and phylogenetic analyses were done using maximum likelihood, neighbor joining and maximum parsimony in MEGA6 (Tamura et al., 2013).

Table (1): Preparation of master mix using Big dye Terminator V3.1 cycle sequencing kit.

| Reagent                    | Amount   |
|----------------------------|----------|
| Big dye terminator v.3.1   | 2µl      |
| Primer                    | 1µl      |
| Template according to quality of band and concentration of DNA | From 1 to 10 µl |
| Deionized water or PCR grade | Complete till to total volume become 20µl |
| Water                      |          |
| Total volume               | 20µl (Mix well, spin briefly) |

5- Antimicrobial sensitivity test
Bacterial inoculums were prepared by overnight broth culture of the isolated bacteria according to Collee et al., (2007). Antibiotic susceptibility testing was carried out by disc diffusion method based on The Clinical and Laboratory Standards Institute (CLSI) guidelines using Muller-Hinton (MH) agar with and without sheep blood. Sensitivity pattern of the isolates were determined against thirteen selected antibiotic discs (sulfamethoxazole-trimethoprim (1.25-23.75μg), chloramphenicol (30μg), tetracycline (30μg), nitrofurantoin (300μg), erythromycin (15μg), penicillin G (10μg), nalidixic acid (30μg), clindamycin (2μg), kanamycin (30μg), cefotaxime (30μg), vancomycin (30μg), amoxicillin (25μg), cefazolin (30μg) (Oxiod). The dimeter of the inhibition zones was measured
according to CLSI guidelines (CLSI Catalogue, 2017)

6- Ethics Statement

All animals were handled according to the regulations of the Animal Ethics Committee at the Faculty of Veterinary Medicine, South Valley University, Qena, Egypt, with good animal practice following the guidelines of Research Code of Ethics (RCEO-SVU) at the South Valley University.

RESULTS

The complication associated with LSD infection were Pneumonia, edema in the brisket, Purulent osselets, nodules on eye lid, nostril, nose, fore head, purulent osselets, rhinitis, udder nodules and fistula sit-fast nodules, ulcer in leg, corneal opacity and endophthalmitis as showing in table (3), Fig (1a,b,c,d and 2e and f).

Table (3) Showing the recorded common microbiological, surgical complications and its isolated in bovine.

| Cases No. | Surgical complication | MOS isolated |
|-----------|-----------------------|--------------|
| 1- LSD    | Pneumonia, Edema in the brisket, Purulent osselets, | Serratia odoriferans |
| 2- LSD    | Nodules on eye lid & nostril and Purulent osselets | Bacillus licheniformis |
| 3- LSD    | Nodule in the nose and forehead, rhinitis, Purulent osselets and udder nodules and fistula | Enterococcus faecium |
| 4- LSD    | Sit-fast nodules, ulcer in leg and corneal opacity | Paenibacillus apiaries |
| 5- LSD    | Nodule in the nose, forehead, rhinitis and endophthalmitis | Microbacterium hominis |
| 6- FMD    | Vesicular lesions, corona band ulcer | Enterococcus hirae |
| 7- FMD    | Swelling of tongue, corona band affection and vesicle on the teat and abortion | Bacillus licheniformis |
| 8- FMD    | Vesicular lesions, swelling of tongue corona band ulcer and blister of the udder | Pseudomonas oxidans |
| 9- FMD    | Swelling of the gum and mastitis and claw over lapping | Trueperella pyogenes |

Figure (1): A: Sit fast nodules all over teat of 3 years old age cow as a complication of L.S.D B: Purulent osselets of 1 year old age bull as a complication of LSD C: Blister of fore limb of 3 year old age cow as a complication of LSD D: Udder nodules as a complication of LSD of 3 years old cow.
Table (4) Showing the results of isolation of micro-organism and results of 16rRNA sequencing.

| Sample | 16rRNA sequencing | Blood agar | Mannitol salt agar | Macconkey agar |
|--------|-------------------|------------|-------------------|-----------------|
| 1      | *Serratia odorifera* | Small grey colony | No growth | Pink colony |
| 2      | *Bacillus licheniformis* | Small brown colony | No growth | no growth |
| 3      | *Enterococcus faecium* | Small white colony | No growth | no growth |
| 4      | *Paenibacillus apiaries* | Long irregular | B- haemolysis | No growth | no growth |
| 5      | *Microbacterium hominis* | White colony | B- haemolysis | No growth | no growth |
| 6      | *Bacillus licheniformis* | Brown irregular | No growth | no growth |
| 7      | *Pseudomonas mossellii* | Small White colony | B- haemolysis | no growth | Pale colour |
| 8      | *Bacillus cereus* | Small grey colony | B- haemolysis | no growth | no growth |
| 9      | *Bacillus licheniformis* | Small White colony | B- haemolysis | no growth | no growth |

The most common complication associated with FMD were coronary band ulcer, vesicular lesions on the gum, tongue and Swelling of tongue and vesicle on the teat and abortion, blister of the udder and mastitis and claw over lapping as showing in (table 3) Fig (2g,h and 3 i, k and l).

The results of conventional isolation and 16sRNA sequencing showing that the most bacterial isolate were isolated from FMD and LSD were *Bacillus licheniformis*, *Enterococcus faecium*, *Paenibacillus apiaries*, *Microbacterium hominis*, *Pseudomonas mossellii*, *Bacillus cereus* and *Trueperella pyogens* table (4), Fig (4), Fig (1S) and Fig (2S). phylogenetic tree was carried by using neighbor joining to compare our isolates with other bacterial isolates in gene bank as showing in Fig (1S). The antibiotic sensitivity tests result for bacterial isolates showing the all bacterial isolates were resistant to more than three antibiotics of different family of antibiotics table (5) and Fig (5).

**DISCUSSION**

The clinical picture of LSD in affected cattle is characterized by skin nodules. The nodules involve the skin and the mucosa of the respiratory tract, eyes, and genital tract. The nodules are characterized by their raised round shape, and pain. Nodules can also develop on the muzzle, buccal mucous membranes, nasal cavity, udder, and teats. Edema develops on the legs and brisket, and enlargement of regional lymph nodes. Suppuration, sloughing, and necrosis are most common complications of the secondary infection of LSD (Ahmed and Dessouki 2013).

Foot-and-mouth disease (FMD) is a devastating and debilitating viral disease of the cloven-hoofed animals and considered a threat to the animal industry globally. (Xu et al., 2013). The FMD is characterized by
high rise of temperature, appearance of vesicles (blisters) on the mouth, muzzle, tongue, snout, nose, teats, inter digital space of feet and other hairless parts of skin (Teifke et al., 2012) which results into off-feeding and lameness. Therefore, direct economic losses due to FMD are attributed to losses in the meat and milk production (Ferrari et al., 2013).

In this study *Serratia odorifera* was isolated from male bull that was suffering from LSD, animal was suffering from pneumonia, coughing, nasal discharge and nodules in mouth and eyes. *Serratia odorifera* species are rod-shaped Gram-negative bacteria, non-spore-forming bacilli that have peritrichous flagella. The organism had a characteristic strong potato-like odor when grew on 5% sheep blood agar (Stock et al., 2002). The normal reservoir of *S. odorifera* is unidentified. Most strains have been isolated from cultivated mushrooms and from clinical specimens (bile, blood, pus and most commonly, sputum) of *S. odorifra* causing an invasive human infection Chmel. H, (1988).

In this study *S. odorifra* are resistant to tetracycline, kanamycin, penicillin, cefotaxime and cefazolin and are sensitive to sulfamethoxazole-trimethoprim, nalidixic acid and amoxicillin. This agrees with Stock et al., (2003) who reported that *S. odorifera* isolates are naturally resistant

Table (5) Showing the results of sensitivity test of ten cases isolated

| Ab                  | Stx | F  | TE | NA | DA | K  | C  | E  | P  | CTX | VA | AX | CZ |
|---------------------|-----|----|----|----|----|----|----|----|----|-----|----|----|----|
| Serratia odorifera  | S   | R  | R  | S  | R  | R  | I  | R  | R  | R   | R  | S  | R  |
| Bacillus licheniformis | S  | R  | S  | R  | S  | R  | R  | R  | S  | R   | R  | I  | R  |
| Enterococcus faecium | R   | R  | I  | R  | I  | R  | I  | R  | R  | R   | R  | R  | R  |
| Paenibacillus apiaries | S  | R  | S  | R  | S  | R  | R  | R  | I  | R   | R  | R  | R  |
| Microbacterium hominis | R  | R  | R  | R  | R  | R  | R  | R  | R  | R   | R  | R  | S  |
| Bacillus licheniformis | S  | I  | S  | R  | S  | R  | R  | R  | R  | R   | R  | R  |
| Pseudomonas mosselii | R   | R  | R  | I  | R  | R  | R  | R  | R  | R   | R  | R  | R  |
| Bacillus cereus      | R   | R  | S  | R  | R  | S  | R  | R  | R  | R   | R  | R  | R  |
| Bacillus licheniformis | S  | S  | S  | R  | S  | R  | R  | R  | R  | S   | R  | R  | R  |
| Trueperella pyogens  | S   | R  | S  | R  | R  | S  | R  | R  | R  | R   | R  | I  | R  |

Ab= antibiotic disc, M.o = Microorganism, Stx = Sulfamethoxazole-trimethoprim
F= Nitrofurantoin, TE= Tetracycline, NA= Nalidixic acid, DA= Clindamycin, K= Kanamycin, C= Chlomphenicol, E=Erythromycin, P= Penicillin, CTX= Cefotaxime, VA= Vancomycin, AX= Amoxicillin and CZ= Cefazolin.

from LSD, animal was suffering from pneumonia, coughing, nasal discharge and nodules in mouth and eyes. *Serratia odorifera* species are rod-shaped Gram-negative bacteria, non-spore-forming bacilli that have peritrichous flagella. The
to tetracyclines and chloramphenicol and are generally susceptible to sulfamethoxazole-trimethoprim, aminoglycosides, ticarcillin, piperacillin, cefotaxime, ceftriaxone, ceftazidime, cefepime, carbapenems, monobactams, and the quinolones.

In this study *Bacillus Licheniformis* was isolated from three cases (one case suffered from LSD and two cases suffered from FMD). The first case was a cow with LSD, which caused multiple nodules on udder and nose and purulent osselets. The second case was a male bull with FMD causing vesicle on the gum and tongue.

The third case was a cow affected with FMD and the clinical signs appeared as coronary band ulcer, swelling and blistering of the tongue. *Bacillus Licheniformis* is Gram-positive bacillus, spores of this bacterium redside in soil, bird feather especially chest, which make it desirable to be used for industrial purpose (Caldow et al., 1996). The presence of *Bacillus Licheniformis* as a complication of LSD and FMD may be attributed to the contamination of animal wounds with dust and soils.

In this study *Bacillus Licheniformis* isolates were sensitive to sulfamethoxazole-trimethoprim and nalidixic acid and resistant to the other antibiotics. The clinical pictures and the isolated microorganisms were nearly similar to that mentioned by Bobkurov, 1977 and frank, 1981. The resistances of *B. licheniformis* to chloramphenicol and streptomycin were intrinsic (Ruegg 2010).

In this study *E. faecium* was isolated from a cow affected with LSD and was suffering from purulent osselets in the fore limb and udder fistula. *Enterococcus faecium* is one of the most important species of the genus Enterococci that includes 38 different species. It is a Gram-positive, facultative anaerobe that causes a variety of infections including, endocarditis, urinary tract infection, intra-abdominal infection and wound infection (Devriese et al., 2006; Murray et al., 2009 and Teixeira et al., 2011).

**Figure (3):** I: Blisters in the mouth causing excess salivation of 2 years old age cow as a complication of FMD; K: Rupture of vesicle of teat of 4 years old age cow as a complication of FMD; L: Blister of the udder of 3 years old cow as a complication of FMD.
This is due to contamination of the wound as a secondary infection. Udder fistula was recorded in this study as a complication of LSD. The isolated bacteria were *Enterococcus faecium* the same result was reported by Abd-El- Hady (1993). In our study *Enterococcus faecium* was intermittently sensitive to tetracycline, clindamycin and chloramphenicol, but was resistant the other antibiotics. In our study the infection was coming from contamination of the wound or as a sequel of infection either by LSD or FMD.

In our study *Paenibacillus apiaries* was isolated from an animal with LSD and was contaminated wound and Sidfast nodules. Animal was located near a honeybee colony, so we suspect that *Paenibacillus apiaries* infection happened due to affection from insects (honeybee colony). *Paenibacillus apiaries* is a Gram-positive bacillus, spore-forming bacterium that can invade honeybee larvae (Spence et al 2013). *Paenibacillus apiaries* isolate was sensitive to sulfamethoxazole-trimethoprim and kanamycin but resistant other antibiotics.

*Microbacterium hominis* was isolated from the eyes of a cow affected with LSD. This is the first case reported in animals and caused endophthalmitis in the cow. *Microbacterium hominis* is a Gram-positive, non-acid fast, non-capsule forming and non-spore-forming bacterium. It is isolated from lung aspiration and causes endophthalmitis in human (Funke et al 1995). The micro-organism was resistant to all antibiotics tested except cefazolin.

*Pseudomonas mosselii* was isolated from a cow affected with FMD. Clinical symptoms included swelling of the tongue and gum and coronary band ulcer. The owner of this animal worked in a hospital; we suspect the infection to be of a nosocomial origin. *Pseudomonas mosselii* is a Gram-negative bacillus that is the third most common cause of nosocomial infection after *Staphylococcus aureus* and *Escherichia coli* (Abbasi et al.,2017). It affects mainly patients with cystic fibrosis, burns or immunodeficiency. The obtained data of antibiotic sensitivity of in our study showed *Pseudomonas mosselii* to be resistant to all antibiotics and this agree with (Khaledi and Meskini,2018).

*Bacillus cereus* was isolated from a cow affected with FMD. Clinically, it showed udder nodules and abortion. *Bacillus cereus* is a Gram-positive bacterium, motile, rod-shaped facultative anaerobic and β-hemolytic bacteria which affects both humans and animals Caldow et al., 1996 stated that the ability of bacillus species to infect the bovine placenta resulting in subsequent abortion, has been studied only for *Bacillus cereus*. The data of antibiotic sensitivity test of *Bacillus cereus* revealed sensitivity to tetracycline but was resistant to all antibiotics used in this study.

Figure (4): PCR results of 16rRNA among ten clinical isolates. Lane L: DNA ladder, Lane p: control positive, Lane N: control negative, lane 1-10 positive for 16 r RNA gene.
Fig (5): Sensitivity test using antibiotic disc. 1 (AX= Amoxicillin), 2 (CZ= Cefazolin), 3 (K= Kanamycin), 4 (VA= Vancomycin), 5 (Cc= Cholmphenicol), 6 (DA= Clindamycin), 7 (CTX= Cefotaxime), 8 (P= Penicillin), 10 (F= Nitrofurantoin), 11 (NA= Nalidixic acid), 12 (TE= Tetracycline) and 13 (STX= Sulfamethoxazole-trimethoprim).

*Trueperella pyogenes* (*T. pyogenes*) is a species of bacteria that are non-motile, facultative anaerobic and Gram-positive. Over the last decades, *T. pyogenes* has been implicated as a cause of different clinical manifestations in domestic animals (Radostits et al 2007 and Greene et al 2012). Quinn et al (2011) stated that bovine intramammary infections caused by *T. pyogenes* are associated with the highest somatic cell count (SCC) in milk and significant losses in milk yield, as well as high percentages of nonfunctional quarters. *T. pyogenes* is the most important bacterial risk factor for clinical mastitis. *T. pyogenes* infections have been usually described to cause pyogenic infections in livestock (Radostits et al. 2007) causing bovine mastitis. In our study the *Trueperella pyogenes* was isolated from a cow affected with FMD which clinically appeared as swelling of the gum, tongue and mastitis and claw affection. The infection may be due to contamination of the udder or occurred as a secondary infection, this agrees with Quinn et al. (2011). Sogstad et al. (2005) Stated that claw overgrowth was highly recorded in dairy cows as a complication of FMD. *T. pyogenes* is a common pathogen that causes suppurative clinical manifestations among cattle and pigs, although it is specially reported as a causative agent of bovine mastitis (Ruegg, 2010). The micro-organism was sensitive to sulfamethoxazole-trimethoprim, tetracycline and kanamycin but was resistant to other tested antibiotics. Diseases caused by *T. pyogenes* generate significant economic losses, mainly in cattle and swine breeding, causing a reduction of meat and milk yield. In addition, it decreases reproductive efficiency and sometimes it is necessary to cull diseased animals. The antimicrobial resistance in *T. pyogenes* is an emerging problem because of the common use of these drugs in agriculture (Feßler and Schwarz, 2017).

In conclusion most bacterial isolates were isolated from LSD and FMD as complication were related to environment and human pathogens.

**ACKNOWLEDGMENTS**

Many thanks to Dr. Mahmoud Hamed Sabra, lecturer of Poultry and rabbit disease Faculty of Veterinary Medicine, South Valley University, for his help in analysis of sequencing data.

**CONFLICT OF INTEREST**

The authors declare that they do not have any conflict of interest.

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