Detection Of The Coa Gene In Staphylococcus aureus From Different Sources By Polymerase Chain Reaction

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ABSTRACT
One hundred and ninety eight samples of human and animal sources from different areas in El-Sharkia Governorate were used in this study to isolate Staphylococcus species and to determine the presence of coagulase (coa) gene, a virulence gene detected in S. aureus (CoPS), by PCR. Primary isolation on mannitol salt agar, β-haemolysis, tube coagulase test (TCT) and other biochemical characterization revealed 32.8% (65/198) of the total samples were infected with staphylococci, only 35 isolates were biotyped as coagulase-positive Staphylococci (Cops) and represented as 8 (8.8%), 10 (22.2%) and 17 (26.9%) in milk, meat and human samples, respectively. Meanwhile, Coagulase-negative Staphylococci (CoNS) were obtained only from cows milk. Fifteen CoPS (5 milk samples, 4 meat products and 6 human samples) were subjected to analysis by PCR for detection of coa gene. The resulted amplicons were 648, 723, 812 and 913 bp. According to coa gene polymorphism, milk and meat isolates were grouped into 5 groups, while human isolates were grouped into 4 groups. It was noted that two of the strains, classified as coagulase negative by tube coagulase test, were found to be positive with PCR amplification of the gene which clearly emphasizes the use of molecular methods in detecting S. aureus. In conclusion, the coa amplification has been considered as a simple and accurate method for typing of S. aureus.

Key words: Mastitis. Coagulase polymorphism. CoPS. CoNS. Food products.

INTRODUCTION
Staphylococcus aureus is considered a significant pathogen of animal and human. Economic importance to dairy industry due to staphylococcal infection results from subclinical mastitis accompanied with reduction in the milk quality and a loss in its production. In spite of increasing
frequency of isolation of coagulase negative staphylococci (CoNS) from the bovine mammary glands, CoPS is recognized worldwide as a major pathogen causing subclinical intramammary infection in dairy cows (Turutoglu et al., 2005). S. aureus is the most common bacteria, followed by E. coli and S. agalactiae in subclinical mastitis cases, while E. coli is the most common bacteria detected in clinical mastitis cases, followed by S. aureus and S. agalactiae (Amin et al., 2011). S. aureus is a predominant colonizer of the skin. It is responsible for many different types of human infections. The predominant type of infection usually related to skin abscesses and can occur in the form of furuncles or carbuncles, but it rarely causes infection without a predisposing route of entry. Impetigo is a well known staphylococcal related infection which is characterized by honey-crusted lesions of the skin (Friedman and Ratard, 2007). S. aureus has a large repertoire of virulence factors, including structural and secreted products that play a role in pathogenesis (Defres et al., 2009). Staphylocoagulase is an extracellular protein that has traditionally been used to differentiate S. aureus from the less virulent staphylococci (CoNS). S. aureus secretes two clotting factors, coagulase (Coa) and von Willebrand factor binding protein (vWbp), the Coa and vWbp together are required for the formation of abscesses and promote the non proteolytic activation of prothrombin and cleavage of fibrinogen, reactions that are inhibited with specific antibody against each of these molecules. Coa and vWbp specific antibodies confer protection against abscess formation and S. aureus lethal bacteraemia, suggesting that coagulases function as protective antigens for a staphylococcal vaccine, so, coagulases may be used as vaccine antigens to elicit antibodies that protect humans against S. aureus infections (Cheng et al., 2010). Moreover, abscesses formation, bacterial persistence in host tissues, blood clotting of coagulase producing S. aureus enable the pathogen to cause lethal sepsis, disseminate as thromboembolic lesion and resist opsonophagocytic clearance by host immune cells. Preclinical evidence suggests that inactivation or neutralization of coagulase may prevent the pathogenesis of staphylococcal infection (Friedman and Ratard, 2007; Chadrakanth et al., 2010 and Mcadow et al., 2012). Rapid and accurate typing of Staphylococcus aureus is crucial to understand the transmission of this infectious organism. The traditional phenotypic methods have several drawbacks (Tenover et al., 1994). Molecular typing can shorten or
prevent an epidemic and reduce the number and cost of nosocomial infections. This typing can also clarify whether the strains from the environment, instruments, staff, or food are responsible for causing infection. In this way this helps to trace the source of infection or an outbreak, tracking the spread of infections and helps to take specific infection control measure (Hacek et al, 1999). The coa gene amplification has been considered a simple and accurate method for typing of S. aureus isolated from distinct sources, the coagulase protein is an important virulence factor of S. aureus. Like spa, coa has a polymorphic repeat region that can be used for differentiating S. aureus isolates. The variable region of coa is comprised of 81bp tandem short sequence repeats (SSRs) (Van-Belkum et al, 1998). The objective of this study was to detect the presence of coa gene in S. aureus from different sources by PCR amplification.

MATERIALS AND METHODS

Samples:

One hundred and ninety eight samples of human and animal sources were collected from different areas in Sharkia Governorate in the period from January to May 2011. Samples of Human source (63) as blood (10), urine (12), C.S.F. (8), sputum (8), pericardial fluid (7), pus (10) and peritoneal fluid (8) were collected from different clinical cases of different ages administered to ultra laboratory and El Gamah hospital. Samples of animal source (135) were mastitic milk (90) collected from 4 farms (Abo-Mandor, Elsalhia, El hosania and Italian company farms) and random meat products (45) as minced meat (20), burger (15) and sausage (10) were collected from different sale markets. All samples were sent as soon as possible to laboratory of bacteriology, Faculty of Veterinary Medicine, Zagazig University and examined for staphylococci.

Isolation and identification of Staphylococcus species:

All samples from different sources were inoculated onto mannitol salt agar (Difco, USA.) and incubated for up to one week at 37°C and all developed colonies with grapes like shape under light microscope were inoculated into soft agar for culture preservation (Difco, USA.). The distribution of the isolates according to growth onto MSA, oxidation fermentation (OF) test and bacitracin susceptibility was detected (Mackie and McCartney, 1996). The developed colonies onto MSA were subjected to tube coagulase test [TCT] (Gillespie, 1943) and for β-haemolysis onto blood agar to differentiate between the CoPS and CoNS. These isolates were further identified as S. aureus by API20S kits, commercial identification
system (BioMerieux, 1992) and kept frozen at -20 °C in nutrient broth with 3% glycerol until molecular tests were carried.

**DNA extraction:**
The bacterial genomic DNA was extracted from only 15 *S. aureus* isolates (CoPS), using DNA extraction kits (Biofermentus).

**DNA amplification:**
The PCR for amplification of *coa* gene was performed in a total reaction volume of 25 µl for one sample according to (Himabindu et al., 2009) using PCR master mix (DreamTaq™ Green Master Mix (2X), Fermentas, Catalogue No. #K1081., Lot: 00055548). It is a ready to use solution containing DreamTaq™ DNA polymerase, optimized DreamTaq™ Green buffer (2X), 4mM MgCl2 and dNTPs (dATP, dCTP, dGTP and dTTP, 0.4 mM each). The sequence of primer used for amplification of *coa* gene (Himabindu et al., 2009) was

forward: 5’CGAGACCAAGATTCAACAG 3’

and Reverse: 5’AAAGAAAAACCACCTCACATC 3’.

The PCR cycling protocol was applied as following: initial denaturation at 94°C for 5 min, followed by 30 cycles of denaturation at 95°C for 30 second, annealing at 55°C for 45 second and extension at 72°C for 2 min., followed by a final extension at 72°C for 7 min.

**Agarose gel electrophoresis and visualization of PCR products:**
5 µl of each amplicon was electrophoresed in 1.5 % agarose gel (Sigma -USA) and visualised under U.V. transilluminator (Spectroline Model TC- 312 A, USA) (Sambrook et al., 1989), then compared with the marker DNA ladder (100 bp, Gene Ruler™, Fermentas).

**RESULTS**

**Culture findings:**
Staphylococci were isolated and identified by traditional phenotypic methods from 32.8% (65/198) of the samples represented 38, 10 and 17 isolates from milk, meat products and human samples, respectively. Only one isolate of micrococcus was obtained from milk samples. On MSA agar, yellow colonies showing mannitol fermentation were selected and divided into 2 groups based on the coagulase test and β-hemolysis.

CoPS were confirmed as golden yellow pigmented colonies, coagulase positive and β-haemolytic 53.8% (35/65) represented 8 (8.8%), 10 (22.2%) and 17 (26.9%) from milk, meat and human samples, respectively.

Meat staphylococci isolates (10) recovered from meat products were distributed as following: (1 from minced meat, 4 from sausages and 5 from burger). Meanwhile, human
staphylococci (17) were 3 from urine, 5 from pus, 3 from sputum, 2 from CSF, one from blood, 2 from pericardial fluid and 1 from peritoneal fluid.

CoPS were predominated in isolates of meat products and human origin (the percentage of coagulase positive was equal to the percentage of β-haemolytic organisms 100%), meanwhile, in staphylococcal isolates of milk samples, the percentage of coagulase positive (21%) (8/38) was less than that of β-haemolytic (26%) (10/38), as two isolates were β-haemolytic but coagulase negative.

CoNS 46.2% (30/65) were isolated only from milk samples and could be confirmed by API 20-S. Biotyping of the isolates by analytical profile index (API20-S) revealed that all coagulase positive and β-haemolytic isolates were S. aureus, in addition to the two isolates of milk samples that were coagulase negative and β-haemolytic.

**PCR findings:**

Most of isolates 80% (12/15) produced a single band, with molecular sizes ranging from 648-913 bp, whereas 2 (13.3%) human isolates yielded 2 amplification products (double amplicons with molecular sizes of 723 + 812bp and 812+913 bp) and only 1 (6.6%) isolate of milk samples yielded 3 amplification products (triple amplicons with molecular sizes of 723+812 + 913 bp) (Photos 1 & 2, Figure 1).

The product 812 bp was the most frequent and accounted for 5/15 (33.3%) of the isolates, followed by 648 bp (26.6%), 723 bp (13.3%) and 913 bp (6.6%). According to coa gene polymorphism, the samples were grouped into 5 groups in (nine isolates of the milk and meat product) and 4 groups in (the six isolates of human subjects) (Figures 2 & 3). The two isolates from milk that were β-haemolytic and negative tube coagulase test, gave specific amplicon by coa gene amplification.

![Fig.1 Genotypic classes of S. aureus by coa gene polymorphism.](image)
Fig. 2 Genotypic classes of *S. aureus* of human origins

![Chart: Genotypic classes of S. aureus of human origins](chart.png)

Fig. 3 Genotypic classes of *S. aureus* of animal origins.

![Chart: Genotypic classes of S. aureus of animal origins](chart.png)

Photo 1: *Cva* gene polymorphism for *S. aureus* isolates of milk and meat products.

DISCUSSION
Molecular typing of microorganisms is now recognized as an essential component of infection control program. These molecular techniques are effective in tracking the spread of nosocomial infections and planning the activities of the infection control program. Production of coagulase is an important phenotypic feature used worldwide for the identification of Staphylococcus aureus (Goh et al., 1992). The coa gene amplification has been considered as a simple and accurate method for typing of Staphylococcus aureus.
In this study, S. aureus was recovered from different human and animal sources, phenotypically and genotypically identified. Seven coa PCR types and 9 groups patterns were detected, which indicated considerable heterogeneity in the coa gene of S. aureus in the studied samples. Until recently, single and also double banded coa PCR products were reported in S. aureus strains derived from bovine mastitis in Brazil (Da Silva and Da Silva, 2005). Double-banded amplification product was detected only in one coa positive isolate (Goh et al., 1992), which explained the presence of double-banded amplification products with different allelic forms of the coa gene.
Our amplification of DNA of S. aureus isolates obtained from human and animal sources revealed four amplicons (723, 812, 648 and 913 bp). These amplicons could classify the isolates into 4 groups (human) or 5 groups (animal).
Similar results were obtained by Himabindu et al. (2009) who showed that the sizes of PCR products obtained after amplification of S. aureus of human subjects rang from 650-1000 bps. Ishino et al. (2007) classified 678 S. aureus isolates of human subjects into 8 classes and the sizes of the PCR products of coa gene ranged from 350 to 917 bp in increments of 81 bp. In addition, Schlegelova et al. (2003) reported the size of coa gene PCR product of S. aureus isolates from dairy cow and human 650 -1050 bp, this result gave 4 classes at 650 ,730 ,810 and 1050 bp ;class 730 bp was the most common class between the isolates. Also Hookey et al. (1999) detected the size of coagulase PCR product of S. aureus isolates from human subjects that was either 660 , 603 or 547 bp. Furthermore, Reinoso et al. (2008) detected that PCR amplification of the coa gene of S. aureus isolated from human, bovine subclinical mastitis and food samples which-yielded seven different coa types from 45 S. aureus strains with amplicon sizes ranging from 400 to 1000 bp. Seven (42%) human infection strains had coa polymorphic regions with an amplicon of 700 bp, indicating the presence of five repeats, Six (40%) of the bovine strains showed an amplicon of 900 bp, indicating the presence of seven repeats and four (50%) strains from food samples showed four repeats.

Also, Da Silva and Da Silva (2005) showed that the amplification of the coa gene from the 64 S. aureus isolated from cow with mastitis produces 27 different PCR-products; which ranged from approximately 579 to approximately 1442 bp. Sizes 790, 759, 725 and 579 bp were the most frequent. Two CoNS isolates showed similarity to S. aureus, and were reclassified as CoPS species by API20s and coa gene detection with 648 and 812 bp amplicons. The same results were obtained by Himabindu et al. (2009) and De Moura et al. (2012) who noted that two of the strains that were classified as coagulase negative by tube coagulase test were found to be positive with PCR amplification of the gene. So the correct amplification of all isolates by PCR not only confirm the results of biochemical tests but is more accurate. Coagulase production is the principal criterion used by the clinical microbiology laboratory for the identification of Staphylococcus aureus. Numerous allelic forms of S. aureus coagulase exist, with each isolate producing one or more of these enzyme variants (Landolo, 1990).

In conclusion, this study has shown that infection in the studied samples was caused by S. aureus strains harboring more than one coa
genotype and that only one genotype predominated. However, further studies using a RFLP technique and nucleotide sequencing methods on a large collection of strains could be conducted to determine the common characteristics of the predominant strains.

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Tenover, F. C., Arbeit R., Archer G., Biddle J., Byrne S., Goering
الملخص العربي

تحديد جين التلزن الدموى في الميكروب العنقودى المعزول من مصادر مختلفة بواسطة تفاعل البلمرة المتوسط

خضعت 198 عينة من مصادر حيوانية وادمية من أماكن مختلفة في محافظة الشرقية لعزل الميكروب العنقودى (المكور العنقودى) وتحديد جين التلزن الدموى به،والذي يعد من جينات الضرورة للمكورات العنقودية موجبة التلزن بواسطة تفاعل البلمرة المتوسط. لقد أثبت تخمير المانيتول والتحلل الدموى واختبار التلزن الإنبوبي بالإضافة إلى بعض الخصائص الحيوية التصنيفية أن 198/65 عينة بنسبة (33.28%) كانت بها عنقوديات، وصنفت 33 عينة منها حيويا كموجبة التلزن الدموى وشملت 8(8.88%) وخلل 117(64.22%)، والتي كانت من عينات الحليب ومنتجات اللحوم والعينات الأدمية على التوالي، علاوة على ذلك تم عزل العنقوديات سالبة التلزن من الحليب فقط.

وخفضت 15 عينة من العنقوديات موجبة التلزن لاختبار تفاعل البلمرة المتوسط لتحديد جين التلزن الدموى بها وقد حصلنا على أربعة أميليونات مفصولة كهربائيًا 248,733,812,913 واستطاعت هذه الأميليونات تصنيف 9 عارات من منتجات الألبان واللحوم إلى 5 مجموعات بينما ال 6 عارات الأدمية إلى 4 مجموعات. لوحظ أن علرات من العنقوديات قد صنفت سالبة التلزن ( بواسطة اختبار التلزن الإنبوبي) لكنها صنفت موجبة التلزن على اساس جين التلزن الموجود بها ( بواسطة اختبار تفاعل البلمرة المتوسط) وهذا يوضح اهمية التصنيف الجيني لجين التلزن كطريقة دقيقة وبسيطة في تحديد ميكروب العنقود الذهبي.