**Incidence and Phenotypic characterization of *Staphyococcus aureus* isolated from mastitic cows.**

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**Abstract**

A total of 412 milk samples were collected from clinically and subclinical mastitic cows (188, 224) respectively and examined bacteriologically for *Staphylococcus aureus* forthe isolation rate and studying of the phenotypic characterization of the isolates. The results revealed that *S.aureus* was isolated in an incidence of 50% and17.5% from clinically mastitic and subclinically mastitic cows respectively. All Staphylococcus isolates showed symmetrically in their phenotypic characterization including cultural characters on different media. Vitek2 system succeeded in providing definitive identification results for gram positive bacterial by identification card (GP) were used for rapid and easy identification of Staphylococcus spp .

**Keywords** :Saureus- clinical-subclinical-mastitis-Vitek2

**1-Introduction :**

Mastitis (inflammation of mammary gland) is one of the most devastating disease conditions leading to significant economic losses globally (Kumar *et al.,* 2010a;

because of reduced milk production, treatment costs, increased labor, milk withholding following treatment, death and premature culling Due to multiple etiologies, it always remained a challenge to veterinarian worldwide. Approximately, 140 species of microorganisms have been identified as etiological agents of bovine mastitis. Of these various etiological agents, *Staphylococcus aureus* is a major pathogen associated with bovine clinical and subclinical mastitis (Tenhagen *et al.,* 2009; Piepers *et al*., 2007; Bhatt *et al*., 2011; Cervinkova *et al.,* 2013).

The mastitis caused by *S. aureus* is characterized by significantly lower cure rates compared with infections caused by other microorganisms, which may be either as a result of unusually frequent acquisition of antibiotic resistance mechanisms among this group of bacteria or also their ability to form biofilm (slime) (Cramton et al., 1999). Considering the potential of the area and the economic significance of dairy production to the local community.

Almost any microbe that can opportunistically invade tissue and cause infection can cause mastitis. About 150 species of microorganisms mostly bacterial is able to cause mastitis.(Wyder *et al*., 2011**)**However, staphylococci, streptococci and other related gram-positive, catalase-negative cocci represent the most important causative agents.(Loonen, *et al* 2012)

Staphylococcus species are aerobically growing Gram-positive cocci . Isolation of Staphylococcus species is usually not difficult since Staphylococci not fastidious organism and will grow on commonly media and under variety of conditions (Rowlinson *et al.,* 2006)

Mastitis is recognized as the most important dairy herd problems worldwide. Economic losses of mastitis include decrease in milk quantity & quality and high cost of treatment *Staphylococcus aureus* is one of the most common etiological pathogens, causing intramammary infections in dairy herds leading to severe economic losses in worldwide industry. Accurate identification of the *Staphylococcus aureus* is therefor of great importance in bacteriological laboratory , The vitek2 system used with gram positive (GP) identification card (biomereux) is an automated machine designed to provide rapid and accurate phenotypic identification for most clinical staphylococcus . (Funk, and Funke, 2005)the present study was carried out to detect the incidence of *Staphylococcus aureus* infection in mastitis cases and their phenotypic characterization .

Recently a new automated identification system such as Vitek2, accompanied by identification cards that give reliable and rapid identification, was internationally reported by Chatzigeorgiou et al.,(2011). In addition, identification of bacteria by VITEK2 system has revealed prominent inter laboratory reproducibility and is quickly being included as a routine method for animal laboratory microbiology .

**2-Material and methods :**

1-Animals

A total of 103 cows were examined in this study were classified into clinical and subclinical mastitic cases as (47) clinically mastitic cows and (56) subclinically mastitic cows.

2-Samples

A total of 412 milk samples of which 188 from clinically infected cases and collected according to**(** Quinn *et al.,* 2002*).*The udder of each animal was palpated before sampling for presence of clinical signs of mastitis, the examined udders were thoroughly washed and carefully dried with clean dry towel. Then the teat was swabbed with 70% ethyl alcohol. After that the first few jets of milk were discarded and milk samples were collected in 50ml sterile falcon tube from clinically affected quarter. As well as 224 milk samples were collected from subclinical infected cows (Normal milk ) after application of California mastitis test (CMT) according to(Rasdostitis *et al* ., 1994

Clinical examination **:** The cases of Clinical Mastitis (CM) were diagnosed on the basis of history, clinical signs, physical examination of udder (swelling and pain) and milk (colour-yellow or blood tinged and consistency-watery, etc), while subclinical mastitis (SCM) was diagnosed on the basis of California Mastitis Test (CMT) (Schalm *et al.,* 1971).

Bacterial isolation and identification: Each of the thoroughly mixed milk sampler(Mastitis/subclinical mastitis) was transferred to 10 mL of nutrient broth and incubated at 37°C for15-18 h to resuscitate the organisms. Thereafter, a loopful of inoculum from the nutrient broth was streaked on to nutrient agar plates and incubated at 37°C for 24 h. Presumptive Staphylococcus colonies (golden/white, round, smooth, glistening, opaque) were picked up and characterized biochemically as Barrow and Feltham (1993).Identification by Vitek2 compact system and gram positive test (GP card) were doneaccording to the manufacture’ sinstruction **(**[Chatzigeorgiou et al., 2011](#_ENREF_12))

**3-Results :**

3.1-Isolation and identification of microbe:

3.1.1 Morphological identification:

Out of 188 milk samples of clinically bovine mastitis 94isolates with incidence (50%) and out of 224 subclinical mastitis 39 isolates with incidence (17.5%) were positive for Staphylococcus

3.1.2-Cultural characteristics:

After aerobic incubation on nutrient agar, mannitol salt agar for 24-48 hours at 37 C°, colonies suspected as Staphylococcus were large, 1- 3 mm in diameter, and well isolated colonies reached4 mm in diameter. The suspected colonies were round, convex, smooth with glistening surface. After aerobic incubation of 33 isolates on Baired-Parker’s agar media for 24-48 hours at 37 C°,10 isolates(30.3%) produced black, shiny, convex colonies with entire margins and clear zone surrounding the colonies with or without an opaque zone. This result confirmed by biochemical identification of 33 suspected Staphylococcus isolates that they were 10 isolates (30.3%)were positive for Catalase test (slide technique), Oxidase test, Oxidation - fermentation of glucose (O-F test), Urease test, Gelatin liquefaction test, Mannitol fermentation test, Coagulase test and showed B-hemolysis on Nutrient agar containing 7.5% NaCl and 5% (V/V) defibrinated sheep blood was used.

**4-Discussion:**

Among 34 samples, 12 (32.29%) showed B-hemolysis on 5% cattle blood agar with circular, small, smooth raised whitish colony. Islam *et al*. (2007b) reported that 89.3% *S. aureus* from bovine origin were hemolytic. This variation was due to the difference in sample origin indicating that raw milk contained less association with *S. aureus* as compared with feces of cattle from where the bacteria were isolated by them**.** After overnight incubation on MS agar media, some plates showed yellow colony and some plates showed whitish colony. All the suspected *S. aureus* which produced B- hemolysis on 5% blood agar were able to ferment mannitol salt agar characterized by the formation of yellow colony and white/transparent colony indicated other *Staphylococcu*s spp., as indicated by Begum *et al.* (2007) and Islam *et al.* (2007a, b).In Gram staining, the organism revealed as Gram positive, violet colored, cocci shaped and arranged in grapes like cluster under light microscope.

In this study, Staphylococcus cultural characteristics on nutrient agar, white, yellow or orange water insoluble pigments were formed. And on mannitol salt agar,yellow or golden yellow water insoluble pigments were formed .Also they were aerobic and facultative, liquefied gelatin and fermented a number of carbohydrates to acid. These results were agreed with Jahan *et al.(*2015).Staphylococcus cultural characteristics on Baired-Parker’s agar media for 24-48 hours at 37C°, produced black, shiny, convex colonies with entire margins and clear zones surrounding the colonies with or without an opaque zone. These results were agreed with that of Roberson *et al.* (1992) and Taylor and Francis (2010).

The incidence of *Staphylococcus aureus* in clinical and subclinical mastitis were shown in Table 1. Overall incidence of *Staphylococcus aureus* in clinical as well as sub clinical mastitis, was 94% isolates out of 188 and 39% out of 224 respectively . The incidence of *Staphylococcus aureus* was higher (50.00%) in clinical mastitis in comparison to that of subclinical mastitis (17.50%) and the incidences of *Staphylococcus aureus* in clinical as well as sub-clinical mastitis were higher. These results are almost in the concurrence of previous study conducted in the region in 2010, which revealed *S. aureus* as a major pathogen in the cases of mastitis in Mathura and its surroundings. The incidence of *S. aureus* was 37.03% and 31.70% in cattle (Kumar *et al.,* 2010a). It clearly indicated the presence of *S. aureus* as most prevailing pathogen in the cases of mastitis in dairy animals. Moreover, it is persisting in the similar pattern not only in clinical cases but also in subclinical cases. Various studies have been conducted in different parts of country to assess the prevalence status of bacterial pathogens in mastitis of dairy animals. Similar to the present findings, Purohit (1990) also reported the staphylococcal mastitis in cows to be 31.94% while Mengistie (2003) , Kivaria *et al.* (2005) and Ranjan *et al*. (2011), reported the incidence to be comparatively as 27.37% in Jharkhand, 27.1% and 21.0%, respectively. However, higher incidence of staphylococcal mastitis was reported by Thennarrasu *et al*. (2003), Wani and Bhatt (2003) and Patel (2007) who reported the incidence of staphylococcal mastitis in cows to be 45%, 44% and 47.06% respectively. The high prevalence of staphylococci has been reported by several researchers (Tuteja, 1999; Kaya *et al.,* 2000) and (Hawari and Dabas, 2008; Tenhagen *et al.,* 2009; Nickerson, 2009; Zutic *et al.,* 2012)**.**All *Staphylococcus aureus* isolates (Table. 1) were found catalase positive, oxidase negative urease positive, failed to grow on Macconkey agar, Voges Proskauer (VP) positive and coagulase positive on being subjected to above mentioned biochemical tests.

Similarly, previous studies conducted by Pandya (1991) and Patel(2007) also reported high percentage positivity of *S. aureus* for coagulase production i.e. 100.00%, where as lower percent positivity of *S. aureus* for coagulase production were also reported earlier by Kato and Kume (1980) 34.50%, Boerlin *et al*. (2003)50.00% and Wani and Bhatt (2003) 51.11%. The presence of 100% coagulase positive isolates in present study further suggests the increase in the number of pathogenic *S. aureus* in dairy animals. This is an alarming condition as in general *S. aureus* are supposed to be non pathogenic commensal organisms.

*Staphylococcus aureus* is the most important bacterial microorganism in bovines causing contagious mastitis and highly economic losses in dairy herds **(**Zecconi *et al.,* 2006).

In the present study bacteriological examination and identification of *Staphylococcus aureus* were depend on gram stain, culturing on Baired parker medium, catalase test, Coagulase tube test and DNase test.

This agreed with (Bedane *et al,* 2012**)** they revealed that *S.aureus* is responsible for about 30% to 40% of all mastitis cases. This high prevalence of *S.aureus* in this study may be explained that transmission of infection occurs during the milking process by milker's hands, contaminated equipments and milking machine (Scherrer *et al,* 2004).

identification by Vitek2 accompanied by Gram positive card is based on established biochemical methods and newly developed substrates measuring carbon source utilization enzymatic activities and resistance. There are 47 biochemical tests and one negative control well; Final results are available in approximately 10 hours or less (Biomerieux user guide, 2006).Our results in agreement with all the authors that found Vitek give reliable, rapid and higher correct identification rate Chatzigeorgiou et al.,(2011).

**5-conclusion:**

The results of the present study clearly showed that *S.aureus* is a major cause of mastitis in dairy farm and its incidence was high in clinical mastitis (50%) than subclinial mastitic cows (17.5%).All *S. aureus* isolates in this study showed that symmetrically of the phenotypic characters . The presence of *S.aureus* in apparently normal milk (subclinical mastitis )cow milk is a potential health hazard. Also that sanitary measures are needed to improve the hygienic conditions during milking

**References :**

Barrow, G.I. and Feltham, R.K.A(1993).Cow an and Steel's Manual for the Identification of Medical Bacteria.3rdEdn.,Cambridge University Press,Cambridge,pp:140-43.

Bedane ,A. ,Kasim,G.,Yohannis,T.,Habtamu, T.,Asseged,B. and Demelash b. (2012):Study on pervelance and risk factors of bovine mastitis in Borona a pastrol and agro pastoral settings of yabella district , Borana zone ,southern Ethiopia America-Ethiopia American –Euraseian, J.Agri.&Environ.Sci.,12(10):1274-1281.

Begum HA, Uddin MS, Islam MJ, Nazir KHMNH, Islam MA, Rahman MT (2007). Detection of biofilm producing coagulase positive Staphylococcus aureusfrom bovine mastitis, their pigment production, hemolytic activity and antibiotic sensitivity pattern. Journal of the Bangladesh Society for Agricultural Science and Technology, 4: 97-100.

Bhatt,V.D.,M.S.Patel,C.G.JoshiandA.Kunjadia,2011.Identification and antibiogram of microbes associated with bovine mastitis. Anim. Biotechnol., 22: 163-169.

Biomerieux (2006):Vitek 2 product information document 510769-4EN1. Biomerieux ,INC.,Durham NC.

Boerlin,P.,Kuhnert, ,P. Hussy, D., and Sehaellibaum, M. (2003).Methods for identification of *Staphylococcus aureus* isolates in cases of bovine mastitis.J.Clin.Microbiol.,41:767-771.

Cervinkova,D.,H.,Vlkova,I.,Borodacova,J.Makovcova and V.Babak *etal*.(,2013).Prevalence of mastitis pathogens in milk from clinically healthy cows. Vet. Med., 58: 567-575. 2013. Prevalence of mastitis pathogens in milk from clinically healthy cows. Vet. Med., 58: 567-575.

 Chatzigeorgiou, K.-S., Sergentanis, T.N., Tsiodras, S., Hamodrakas, S.J., Bagos, P.G., 2011. Phoenix 100 versus Vitek 2 in the identification of gram-positive and gram-negative bacteria: a comprehensive meta-analysis. Journal of clinical microbiology49, 3284-3291.

Cramton,S.E.,C.Gerke,N.F.Schnell,W.W.NicholsandF.Gotz,1999.The intercellular adhesion (ica) locus in Staphylococcus aureus and is required for biofilm formation. Infect. Immun., 67: 5427-5433.

De FreitasGuimaraes F., Nobrega, D.B., Richini- Pereira, V.B., Marson, P.M., FigueiredoPantoja, J.C. and Langoni, H. 2013. Enterotoxin genes in coagulase- negative and coagulase-positive staphylococci isolated from bovine milk. J. Dairy Sci. 96:2866-2872.

Funke, G. andFunke,K. (2005).Performance of the New VITEK 2 GP Card for Identification of Medically Relevant Gram-Positive Cocci in a Routine Clinical Laboratory. *J. Clin. Microbiol.*, 43(1):84-88.

Hawari, A.D.andF.Al-Dabbas(,2008).Prevalence and distribution o f mastitis pathogens and their resistance against antimicrobial agents in dairy cows in Jordan .Am.J. Anim.Vet.Sci.,3:36-39.

Islam MJ, Uddin MS, Islam MA, Nazir KHMNH, Rahman MT, Alam MM (2007b). Detection and characterization of coagulase-positive *Staphylococcus aureus*of bovine origin producing enterotoxins and toxic shock syndrome toxin-1. The Bangladesh Veterinarians, 24: 27-33.

Islam MJ, Uddin MS, Nasrin MS, Nazir KHMNH, Rahman MT, Alam MM (2007a).Prevalence ofenterotoxigenic and toxic shock syndrome toxin-1 producing coagulase positive *Staphylococcus aureus*in human and their characterization. Bangladesh Journal of Veterinary Medicine, 5: 115-119.

Jahan M., Rahman M., Parvej M.S., Chowdhury S.M.Z.H, Haque ME, Talukder, M.A.K, Ahmed, S. (2015).Isolation and characterization of *Staphylococcus aureus*from raw cow milk in Bangladesh. Journal of Advanced Veterinary and Animal Research, 2: 49-55

Kaya,O., Kirkan, S. , Gulal M., andUnal, B . (2000).Identification and antibiotic susceptibility of microbes causing mastitis in dairy cows.Vet.Bull.,70:290-290.

Kato, E. and Kume, T. (1980). Enterotoxigenicity of bovine *Staphylococci* isolatedfrom California mastitis test-positive milk in Japan.Japanese J.Vet.Res.,28:75-85.

Kumar, A., Rahal, A. , Dwivedi, S.K. and.Gupta, M.K (2010a).Bacterial Prevalence and Antibiotic Resistance Profile from Bovine Mastitis in Mathura, India..J.DairySci.,38:31-34.

Kivaria,F.M., . Noordhuizen, J.P.T.M Kapaga A.M. and Hogeveen, H. (2005).Risk indicators associated with *Staphylococcus aureus* subclinical mastitis in small holder dairy cows in Tanzania. Proceeding s of the4th IDF International Mastitis Conference,June12-15,2005, Maastricht ,The Netherlands,pp:722-727.

Loonen, A., Jansz, A., Bergland , J., and Wolffs , P., (2012): Comparative study using phenotypic ,genotypic and proteomic methods for identification of coagulase negative staphylococci . J.Clin.Microbiol.,50:1437-1439.

Mengistie,A.Z.,(2003).Molecular Epidemiology of *Staphylococcus aureus* and Streptococcus Agalactiae Isolated from Bovine Mastitis in Ethiopia. Mensch and Buch- Verlag,USA., ISBN:9783898205122,Pages:139.

Nickerson,S.C.,(2009). Control of heifer mastitis:Antimicrobial treatment-an overview.Vet.Microbiol.,134:128-135.

Piepers, S., L. De Meulemeester, A. de Kruif, G., Opsomer, H.W. Barkema and S. de Vliegher, (2007).Prevalence and distribution of mastitis pathogens in subclinically infected dairy cows inFlanders, Belgium. J. Dairy Res., 74: 478-483.

Pandya,K.,(1991).Incidence of *Staphylococcus aureus* in cow milk and assessment of characteristics associated with its virulence .M.Sc. Thesis, Gujarat Agricultural University, Sardarkrushinagar.

Patel,N.P.,(2007).Determination of virulence factors in *Staphylococcus aureus* isolated from clinical cases of mastitis in sheep, goats, cattle and buffaloes. M.Sc.Thesis, Sardarkrushinagar Dantiwada Agricultural University, Sardarkrushinagar.

Purohit, J.H., (1990).Isolation and characteristics of *Staphylococcus aureus* from bovine milk.Ph.D.Thesis, Gujarat AgriculturalUniversity

Quinn, P.J.; Carter, M.E. and Markey, B.K. (1994).Mastitis. In: Quinn, P.J. (ed). Clinical Veterinary Microbiology..Wolfe, Baltimore. pp.327-344

Quinn, P.J., Makey, B.K., Carter, M.E., Donnelly, W.J. and Leonard, F.C. (2002).Veterinary Microbiology and Microbial Diseases. Blackwell Science Ltd.

Ranjan, R.,Gupta M.K andSingh, K.K. (2011).Study of bovine mastitis in different climatic conditions in Jharkh and,India.Vet.World,4:205-208.

Rasositis,O.M., Blood, D.C.,Gay, C.C.(1994):Veterinary Medicine 8thed.ElBSBailliereTiendall London.,Sardarkrushinagar.

Roberson, J. R., Fox, L.K., Hancock, D.D. and Besser, T.E. (1992). Evaluation of methods for the differentiation of coagulase-positive staphylococci. J. Clin. Microbiol.,30:3217-3219.

Rowlinson,M.C.,LeBourgeois,P.,Ward,K.,Song,Y.,Finegold,S.M.,Bricker,D.A.(2006):isolation of strictly anerobic strains of staphylococcus epidermidis.J.Clin.Microbial.,44(3):857-860.

Schalm, O.W., Carroll, G., and.Jain, N.C (1971).Bovine Mastitis. Lea and Fibiger, Philadelphia, USA.,Pages:360.

Scherrer*, D.;*Coti,S.; Muehlberr,J.E.; Zweife ,C. and Stephan , R. (2004):Phenotypic and genotypic characteristics of *S. aureus* isolates from raw bulk – tank milk samples . Veterinary Microbiology .101-101-107.

Taylor and Francis Group, LLC.( 2010). International Standard Book Printed in the United States of America Number-13: 978-1-4398-0408-7.

Tenhagen, B.A., I. Hansen, Reinecke A., and Heuwieser, W. ( 2009). Prevalence of pathogens in milk samples of dairy cows with clinical mastitis and in heifers at first parturition. J. Dairy Res.,76: 179-187.

Thennarrasu, A.,M.R .Muralidharan andMurugan, M. (2003). Incidence o f clinical mastitis in bovines-a study in Chennaicity. Cherion,32:140-141.

Tuteja,F.C.,(1999).Studies on mastitis in buffaloes with reference to serum selenium status and control by treating teat canal infections. Ph.D. Thesis ,Chaudhary Charan Singh Haryana Agricultura lUniversity ,Hisar, Haryana.

 Wani,S.and.Bhatt, M.(2003).An epidemiological study on bovine mastitis in Kashmir valley.IndianVet.J.,80:841-844.

Wyder AB, Boss R, Naskova J, Kaufmann T, Steiner A, Graber HU (2011): Streptococcus **spp. and related bacteria: their identification and their pathogenic potential for chronic mastitis – a molecular approach.**Res Vet .Sci ., **91:**349-357.

Zecconi , A.; Calvinho, L.F. and Fox , K.L.(2006): Staphylococcus aureus intrammamry infections. IDF Bulletin, 408.1-42.

Zutic,M., Cirkovic, I.,Pavlovic ,L,.Zutic, J., Asanin, J.,Radanovic O., andPavlovic, N. (2012).Occurrence of methicillin-resistant *Staphylococcus aureus* in milk samples from Serbian cows with subclinical mastitis. Afr. J.Microbiol.Res.,6:5887-5889.

 **Table 1: incidence of *Staphylococcus aureus* in the mastitis cows**

Animal case no. no. of quarter *S.aureus* isolates

 No %

|  |
| --- |
| Clinically mastitic 47 188 94 50 Subclinical mastitic 56 224 39 17.5Total 103 412 133 30.9 |

**Identification of *Staphylococcus aureus* using VITEK2 compact system**

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 Identification information Card: GP Lot number:242381940 Expires:May29,201713:CDT

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 Completed Apr 19,2016 17:40CDT Status :final Analysis Time:4.75 hours

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Selected Organism 99%probability *Staphylococcus aureus*

Bionumber:10402062763231 Confidence: Excellent identification

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