New Method for Early Detection of two Random Amplified Polymorphic DNA (RAPD) Groups of *Staphylococcus aureus* Causing Bovine Mastitis Infection in Paraná State, Brazil

Dicezar Gonçalves1,2*, Jane Eire Gabriell, Humberto Maciel França Madeira, Guilherme Schnell e Schühli and Vânia Aparecida Vicente

1Divisão de Bioprocessos e Biotecnologia; Universidade Federal do Paraná; 81531-990; Curitiba - PR - Brasil.
2Pontifícia Universidade Católica do Paraná; Curitiba - PR - Brasil. 3Universidade Estadual de Ponta Grossa Ponta Grossa - PR - Brasil

**ABSTRACT**

The aim of this work was to develop a fast and accurate molecular approach to allow early detection of two RAPD groups of *S. aureus* causing bovine mastitis. Seventy five *S. aureus* isolates from infected animals were characterized by RAPD. Genomic fragments isolated from the unique bands present in either group were cloned and sequenced. Based on the DNA sequences, specific primers were designed to allow for the simultaneous detection of either group by multiplex PCR of *S. aureus* DNA isolated from clinical and subclinical bovine mastitis. Results showed that these proposed primers set could be used to detect various clinical and subclinical *S. aureus* isolates as well as the detection of the microorganism in bulk milk. Their use as a specific method for effective and early diagnostic tool for *S. aureus* infection in dairy herds is suggested.

**Key words:** *Staphylococcus aureus*, molecular markers; subclinical mastitis, clinical mastitis, mastitis detection

**INTRODUCTION**

The economic importance of the *S. aureus* causing clinical and subclinical bovine mastitis is largely recognized (Myllys et al, 1997). Clinical mastitis is the single most costly disease of dairy cattle resulting in the reduction of milk yield and quality (Riekerink et al, 2007a). The control of precalving subclinical mastitis by somatic cell count (SCC) is an indicator of intramammary infection (IMI) in bovines (Riekerink et al, 2007b). Udder edema and number of neutrophils in milk (Detilleux, 2004) are also important factors (Compton et al, 2007a). Prevalence of subclinical mastitis was diagnosed in 18.5% of quarters of the udder (Compton et al, 2007b) and increase in standard deviation log of SCC was associated with the presence of *S. aureus* clinical mastitis in bovines (Green et al,(2004). Mastitis persists for long-term and may be detected by Pulsed Field Gel Electrophoresis (Rabello et al, 2005; Anderson and Lyman, 2006). *S. aureus* was isolated coincidently from the herd with problems of increased number of milk SCC, neutrophils and decreased milk production in infected animals. Studies by Cullor (1992) and Gonçalves and Kozicki, (1997) indicated that the disease was present in 15% of all the dairy herds in the world, with a prevalence rate of 50% in
U.S.A. and classified as either contagious or environmental based upon their primary reservoir and transmission mode (Parker et al, 2007). 

*S. aureus* is a contagious pathogen commonly transmitted among the cows by contact with infected milk and the infection reach up to 32% of the herd (Pitkälä et al, 2004). This pathogen is particularly important because it causes mainly subclinical forms of infectious mastitis that are often difficult to detect by the herdsman (Compton et al, 2007b). The evolution of the infection turns difficult its treatment and increases the disease level (Reksen et al, 2006). Upon infection of the secretory tissue, the pathogen diffuses and encysts (Smith, 1977 and Hensen et al, 2000). This condition reinforces the necessity of early diagnosis to prevent the spread of the disease and management of the contaminated milk with *S. aureus*.

In Paraná State, Brazil, the estimated annual economic loss due to mastitis range from $95.62 to 142.42 per cow (Swinkels et al, 2005). These costs include reduced yield, discarded milk, drug therapy, veterinarian costs, premature culling, and increased labor. Fat and protein production is reduced by 9.0% in the clinical mastitis cases (Hagnestam et al, 2007).

Several biochemical and molecular methods are used in the investigation of *S. aureus* in bovine infections (Hartstein et al, 1989; Williams et al, 1990; Prevost et al, 1991; Kosteman et al, 1992 and Kapur et al, 1995). Current identification methods are based on microbiological culture of milk and biochemical tests on the isolated bacteria. At present, species identification by standard methods is laborious and takes at least 2 to 3 days to yield results. Other tests are commonly employed in Brazil to detect the occurrence of the subclinical mastitis, as California Mastitis Test, (Schalm and Noorlander, 1957; Riekerink et al, 2007b). However, since specific identification of the causal agent in Brazil is limited to tradicional methods, PCR-based approaches have been developed to identify the mastitis pathogens, which provides a promising alternative for the rapid identification of bacteria. In addition, the sensitivity of PCR-based assays tends to be better than bacterial cultures (Forsman et al, 1997; Kim et al, 2001) allowing the detection of small number of microorganisms.

These methods may be extremely important when rapid and accurate identification of pathogenic bacteria is required. Different PCR-based methods have been developed for specific and sensitive detection of mastitis pathogens directly in milk. Forsman et al, (1997) identified by PCR test the *Staphylococcus* spp. isolates in Russia, Germany and Denmark, but no cross reactivity could be observed investigating staphylococcal reference strains. Riffon et al, (2001) developed molecular probes reacting in PCR with bacterial DNA from bovine milk providing direct and rapid detection of *Escherichia coli*, *S. aureus*, *Streptococcus agalactiae*, *S. dysgalactiae*, *S. parauberis* and *S. iberis*, using universal primers in Canada. In this bacterial group, Meiri-Bendek et al, (2002) using 16S subunit of the rRNA genes, developed a specific PCR reaction to detect *Streptococcus agalactiae* in the milk of subclinical mastitis. Cremonesi et al, (2006) developed a method to extract DNA directly from the main Gram-positive bacteria known to cause bovine mastitis. Some alternative methods for diagnosis were also developed for detection in cows. Puetkes et al, (2001) and Eckersall et al, (2006) developed a multiplex PCR approach to detect the disease directly from milk cows or alternatively from dairy products, using mRNA 23S and genomic DNA. Youngerman et al, (2004) using CXCR2, a chemokine receptor required for neutrophil migration to infections sites and which contains single nucleotide polymorphisms (SNP) within the gene, showed association of CXCR2 genotypes, with subclinical and clinical mastitis. Ramesh et al, (2002) and Graber et al, (2007) obtained high correlation (*R* = 0.925), with the PCR methods to diagnose *S. aureus* mastitis. Gillespie and Oliver, (2005), using RT-PCR identified 91.7% of *S. aureus* in milk and Moon et al, (2007), based on coagulase gene polymorphism, observed relations with enterotoxins-producing by *S. aureus* strains.

The objective of this work was to detect *S. aureus* by means of multiplex PCR using a set of specific primers, by previous extraction of DNA, cloning and sequencing of monomorphic bands from isolated *S. aureus* DNA in order to make a rapid and specific identification of the microorganism.

**MATERIALS AND METHODS**

A total of 75 *S. aureus* isolates were isolated from the bovine udder proceeding from 240 different cases of clinical and subclinical infections. Morphological and biochemical identifications were carried out following Koneman et al, (2001).
Samples were stored in 7.5% NaCl agar manitol at 4°C. They are cultivated in BHI broth at 37ºC for 43 h. Genomic DNA was isolated by means of CTAB (cetyltrimethylammonium bromide). (Dellaporta et al. 1983; Doyle and Doyle 1987; Daneshwar and Sher-Ullah, 2004).

Genetical variability and polymorphism analysis of 75 isolates of *S. aureus*

To carry out the Random Amplified Polymorphic DNA (RAPD) technique the following primers were used: OPJ 5 (5’CTC CAT GGG G 3’); OPJ 6 (5’ TCG TTC CGC A 3’); Primer 786 (5’ GCG ATC CCC A 3’); Primer 797 (5’ AGC GTC ACT G 3’) and Primer 798 (5’ TGA CCC GCC 3’), based in Myllys et al, (1997) and Pereira et al, (2002) and OPA 2 (5’ TGC CGA GCT G 3’) (Spolidorio et al, 2003). The DNA amplifications reactions were made according Pereira et al, (2002). A final volume of 14.9 µL contained 20ng of purified DNA and 1.0 U of *Taq* DNA polymerase, 3 mM 1 M 1 M of MgCl2, 50 mM 1 KCl, 10 mM 1 Tris-HCl (pH 8.8), 200 µM of Mix dNTP, and 0.5 mM of each primer. The mixture was subjected to a initial denaturation at 94 ºC for 5 min, followed by 45 cycles of 30 s at 94°C, 30 s at 36°C and 1 min 30 s at 72°C. A subsequent final cycle of extension at 72°C for 1 min was carried out in a thermocycler (PT100 Perkin Elmer). The amplification products were analyzed in 1.6% agarose gel (TBE buffer at 114 mV). Two strains spp. one out-group were also included (Dellaporta et al, 1993; Hillis and Bull, 1993).

Isolation of the RAPD genomic DNA fragments

Two polymorphic and monomorphic bands were selected, corresponding to both mastitis cases types and the isolation of DNA fragments was due to the similarity in each group. The DNA fragments were cut and purified by the Purelink Quick Gel Extraction Kit (Invitrogen Life).

**Fragment cloning in *E. coli***

Fragments were prepared according to the pGEM® Systems, (Promega Corporation) and cloning in the *E. coli* TOP10 F1 (Soares et al, 2001). For visualization of the possible insertion fragments in the plasmids, electrophoresis in agarose gel (0.8 %) was carried out after the treatment with EcoRI restriction enzyme (Invitrogen Life).

**AmpliSeq reactions**

All the reactions were prepared with the Kit DYEframe™ ET dye terminator MegaBACE™ (Amersham). Universal primers used in the reactions for plasmid amplification were synthesized and purified by means of Invitrogen™ Life, according Murphy et al, (2005): pUC/M 13 Forward Sequencing Primer: 5’ CGT TGT TGT AAA ACG ACG GCA AGT 3’ and pUC/M13 Reverse Sequencing Primer: 5’ TTT CAC ACA GGA AAC AGC TAT GAC 3’. Reaction systems consisted of 3.0 µL mix ET dye terminator; two primers in equal concentration 5-500 mol 1⁻¹; dNTPs in a final concentration of 125 mol 1⁻¹, 50-500 mg 1⁻¹ of DNA in a final vol. of 7.5µL. All the reactions were cycled at 95°C for 2 min, followed by 35 cycles at 95°C for 20 s and 60°C for 2 min. The PCR product was purified by ammonium acetate 7.5 mol 1⁻¹ and precipitated by ethanol in successive centrifugations (Murphy et al, 2005). Samples were sequenced by electrophoresis in an ABI 377 (Applied Biosystems). The primers were designed considering the sequences obtained.

**Sequences edition and primer design**

Edited sequences (GenBank: Access AM749799 to *S. aureus* RAPD clinic fragment and AM749800 to *S. aureus* RAPD subclinical fragment), examined trough PerlPrimer (Marshall, 2004) allowed the design of the specific primers set for clinical and subclinical strains. For the subclinical strain, the forward primer FSBSTAPH: 5’ CCA AAC TTG GTA ATT GAA GGA C 3’, displayed an optimum annealing temperature of 59.78°C and the reverse primer RSBSTAPH: 5’ ACA ATG TTG AAA CTC AGA CGT C 3’, displayed an optimum annealing temperature of 61.41°C. For the clinical strain the designed primers were the forward FCSTAPH: 5’ TTG ATC TGG ATA CGA TGG AAA GG 3’ with an optimum annealing temperature of 62.48°C and a reverse primer RCSTAPH: 5’ TTA AGT ATA.
AGC TTC ACG CCG AG 3’ with an optimum annealing temperature of 62.72 ºC.

Markers evaluation and protocol development
A multiplex PCR according to Vandercan et al (1995), Pinekter et al (2001) and Vannuffel et al (2003) was prepared applying the proper annealing temperature suggested by the software (PerlPrimer) with a final volume of 25.0 µL. The reaction comprised 2.5 µL of PCR buffer minus Mg (final concentration 1X), 1.0 µL of mix dNTP’s (final concentration of 0.2 mM), Primer mix in final concentration of 0.5 µM, 2.5U of Taq, 25.0 ng of genomic DNA (0.5 µl) and 17.5 µL of Milli Q filtered water. The multiplex PCR products were examined with agarose gel 1%.

RESULTS AND DISCUSSION

Strain recognition and DNA isolation
In the present work, 31.25% of the mastitis cases sampled were confirmed as S. aureus infection by morphological and biochemical analysis (Gonçalves et al 2001). Several procedures for the early identification and fast recognition of S. aureus infections are essential for health studies (both animal and human) due to economical importance in dairy industry. Amplification analysis allowed the identification of two main groups (Fig.1) based on the presence of distinct monomorphic bands. These bands were fully congruent with the identification of the clinic and subclinical strains of S. aureus.

A multiplex PCR according to Vandercan et al. (1995), Pinekter et al, (2001) and Vannuffel et al, (2003) was prepared applying the proper annealing temperature suggested by the software (PerlPrimer) with a final volume of 25.0 µL. The reaction comprised 2.5 µL of PCR buffer minus Mg (final concentration 1X), 1.0 µL of mix dNTP’s (final concentration of 0.2 mM), Primer mix in final concentration of 0.5 µM, 2.5U of Taq, 25.0 ng of genomic DNA (0.5 µl) and 17.5 µL of Milli Q filtered water. The multiplex PCR products were examined with agarose gel 1%.

In addition to its fast performance, the RAPD method assures a high confidence in its diagnosis, being very sensitive to S. aureus studied strains. RAPD data allowed a clear distinction between two clades in the identification of the strains (Fig.1). The identification was fully congruent with the clinical picture noted in the studied infections. Each group of strains might correspond to one cluster of mutually similar genotypes.

![Figure1](https://example.com/figure1.png) - Some of the strains studied show the formation of two S. aureus groups (I - Clinical and II - Subclinical) with low similarity between the groups but with very high levels of security (p< 0.001).
AmpliSeq reactions
Both inserts were effectively amplified with the proper PCR reaction suggested by the plasmid manufacturer protocol (pGEM® Vector Systems). The results are presented in Fig 3. The two distinctive monomorphic bands were recovered displaying the same size of the original excised bands (950 bp for the subclinical sample and 800 bp for the clinical) (Fig.3).

Multiplex PCR
Amplification of the desired regions starting from genomic DNA was always successful using the four designed primers. It was equally effective for single PCR or multiplex PCR reactions carried out under similar conditions. Figure 4 shows agarose gel (1%) for the specific identification of clinical and subclinical isolated strains after multiplex PCR.

Figure 2 - Size and intensity (in maximum absorbance) by samples proceeding from clinical and subclinical mastitis. Bands were eluted with Purelink kit. A star indicates the position of band peak of interest (around of 950 for the subclinical sample and 800 for the clinical).

Figure 3 - Positively cloned strains in agarose gel (1%). Ld = ladder (1 kbases). The 3 kbases bands correspond to the plasmid pGEM®-T Easy Vector previously digested by EcoRI.
In this study, 75 field *S. aureus* were identified using the set of four primers resulting in amplifying the desired fragment and the reference strains ATCC 25923 and ATCC 6538 samples. Also, *S. aureus* in milk samples collected directly from the milk bulk tank according to Kim et al., (2001), (for total DNA extraction) were positively detected. To assure the specificity of the primers, the amplification of *S. mutans* and *E. coli* was attempted but the primers failed in their amplification. The development of a fast diagnosis tool of the subclinical strain was priority. The multiplex PCR approach allowed the detection of both types of strains. The method was able to detect successfully *S. aureus* contamination even in high-diluted milk samples as the ones taken from a 5000L bulk milk tank. These results suggested another useful application to detect eventual contaminations in the stored milk. This protocol could be used as a quality inspection tool for producers and in the chain of distribution, contributing to healthy herds and high ranks of quality for the milk producers.

ACKNOWLEDGMENTS

The authors are grateful to Dr. Emanuel M. de Souza, Mr. Helisson Faoro and Mr. Valter Baura (Biochemistry Department to UFPR) for their technical assistance. Thanks are due to PUCPR and UFPR, for the financial support.

**RESUMO**

Esta pesquisa objetivou o desenvolvimento de técnica rápida e eficiente para diagnosticar precocemente diferentes linhagens de *S. aureus* causadoras de mastite bovina. Como resultados da metodologia empregada, foram isoladas duas linhagens destas bactérias que causam diferentes tipos de mastite bovina. Os fragmentos de DNA genômico caracterizando ambas as linhagens, por meio de RAPD foram inseridos em vetor plasmidial *pGEM* e clonados por meio de clones T10 F1 de *Escherichia coli*. As sequências obtidas permitiram desenhar iniciadores específicos para o reconhecimento de ambas as linhagens, os quais foram testados com amostras de *S. aureus* e com outras linhagens próximas. O diagnóstico por meios moleculares, pode ser realizado diretamente de amostras coletadas de rebanhos leiteiros assim como dos equipamentos de ordenha. A significância deste estudo consiste em um rápido e acurado método para localizar animais infectados, representando importante ferramenta no manejo do rebanho, na redução de custos com tratamentos e, rápida recuperação de rebanhos infectados.
REFERENCES


Marshall, O.J. (2004), PrimerPrimer: cross-platform, graphical primer design for standard, bisulphite and real-time PCR. Bioinformatics, 20, 2471-2472


Received: May 13, 2008; Revised: October 21, 2009; Accepted: October 21, 2009.