Short Paper

Enterotoxin gene profiles among *Staphylococcus aureus* isolated from raw milk

Nazari, R.^{1*}; Godarzi, H.¹; Rahimi Baghi, F.¹ and Moeinrad, M.²

¹Department of Microbiology, College of Science, Qom Branch, Islamic Azad University, Qom, Iran; ²Department of Microbiology, College of Science, Zanjan Branch, Islamic Azad University, Zanjan, Iran

*Correspondence: R. Nazari, Department of Microbiology, College of Science, Qom Branch, Islamic Azad University, Qom, Iran. Email: Nazari1102002@yahoo.com

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Summary

Milk is considered a nutritious food because it contains several important nutrients including proteins and vitamins. Conversely, it can be a vehicle for several pathogenic bacteria such as *Staphylococcus aureus*. This study aimed to analyze the frequency of genes encoding the nine Staphylococcal enterotoxins (SEs) and enterotoxin gene profiles in *S. aureus* isolates derived from raw bovine milk. A total of 52 *S. aureus* isolates were obtained from 246 milk samples of 246 dairy cows from eight different farms in Qom, Iran. On the basis of cultural and biochemical properties as well as by amplification of the 23S rRNA specific to *S. aureus*, all isolates could be identified as *S. aureus*. Of the 52 isolates studied, 80.7% were positive for one or more genes encoding the enterotoxins, and 12 different genotypes were identified. The gene encoding for enterotoxin A (*Sea*) was the most frequent (16 isolates, 30.7%), followed by *Seb* (14 isolates, 26.9%) and *Sed* (8 isolates, 15.37%). Among the genes encoding the other enterotoxins, *Seg* and *Seh* were the most frequently observed (8 isolates each, 15.38%), followed by *Sej* (6 isolates, 11.5%) and *Sei* (1 isolates, 3.84%). With the recent identification of new SEs, the frequency of enterotoxing has increased, suggesting that the pathogenic potential of Staphylococci may be higher than previously thought. These results of enterotoxin genes positivity of milk-derived Staphylococci constitute a potential risk for consumers' health.

Key words: Staphylococcus aureus, Staphylococcal enterotoxins, Raw milk, PCR

Introduction

Staphylococcus aureus is a gram-positive bacterium, which produces many important virulence factors including Staphylococcal enterotoxins (SEs) which are responsible for Staphylococcal food poisoning (SFP), a major type of foodborne illness (Balaban and Rasooly, 2000). Classical SEs have been divided into five serological types (SEA through SEE) on the basis of their antigenicities. In recent years, the existence of new types of SEs (SEG-SEU, except SES and SET) have been reported (Ren et al., 1994; Su et al., 1995; Munson et al., 1998; Zhang et al., 1998; Jarraud et al., 2001; Orwin et al., 2001). Staphylococcal enterotoxins are resistant to inactivation by gastrointestinal proteases such as pepsin. In addition, they displayed strong thermo resistance, for example, SEA retains some biological activity after 28 min at 121°C (Anderson et al., 1996). Staphylococcal enterotoxins can be routinely detected by immunoassay, e.g. enzyme linked immunosorbent assay (ELISA), immunodiffusion, radioimmuno-assay and latex agglutination but the availability of these methods are usually limited to commercial tests for classical SEs (Omoe et al., 2002). Therefore, the DNA-based approach (PCR assays) is thought to be an essential tool for investigating SE genes (Omoe et al., 2002). The identification of S. aureus at the species level is based on amplification of target genes highly conserved within the species. In addition to the 16S rRNA gene, the well established standard target for the identification of bacterial species (Amann *et al.*, 1995), the 23S rRNA genes has proven useful for identification of *S. aureus* at the species level (Akineden *et al.*, 2001; Phuektes *et al.*, 2003). The aim of this study was to identify *S. aureus* strains via amplification of the genes encoding the 23S rRNA and to analyze the genes encoding the Staphylococcal enterotoxins SEA, SEB, SEC, SED, SEE, SEG, SEH, SEI and SEJ in *S. aureus* strains isolated from raw bovine milk.

Materials and Methods

Sample collection and identification

A total of 52 *S. aureus* isolates were collected from milk sample of 246 cows from eight different farms in Qom, Iran. The identification of the isolates was performed by gram staining, catalase, hemolysis and tube coagulase test (Liofilchem, Italy) and was confirmed by PCR amplification of species specific parts of the *S. aureus* 23S rRNA gene.

Genotypic characterization

The DNA of isolates was prepared with the high pure

PCR template preparation kit (Roche, Germany) as described by the manufacturer. The sequences of the oligonucleotide primers, the predicted PCR product sizes and the references are summarized in Table 1. For PCR amplification, the reaction mixture (25 µl) contained 15 ng DNA template, 1 µl of primer F (10 pmol), 1 µl of primer R (10 pmol), 2.5 µl 10 X PCR buffer (Fermentas, Lithuania), 0.5 µl dNTP (10 mM, Fermantas, Lithuania), 1.5 µl Mgcl₂ (25 mM, Fermentas, Lithuania), 0.5 U Taq DNA polymerase (Fermentas, Lithuania) and doubledistilled water to the final volume of 25 µl. DNA amplification was performed in a thermal cycler (Eppendorf, Germany) with initial denaturation at 94°C for 5 min followed by 35 cycles of denaturation at 94°C for 1 min, primer annealing 58°C for 1 min and extension at 72°C for 1 min, followed by a final extension at 72°C for 7 min. The amplified PCR products were electrophoresed in a 1% agarose gel (Fermentas, Lithuania) containing ethidium bromide (Fermentas, Lithuania) and visualized by trans illumination under UV. Molecular size marker (Vivantis, Malaysia) was included in each agarose gel. The S. aureus reference strains ATCC 19095, ATCC 23235, ATCC 14458, ATCC 700699, ATCC 27664, ATCC 25923 were used as positive controls. The reference strains Staphylococcus epidermidis ATCC 12228, Staphylococcus saprophyticus ATCC 14448 were used as negative controls.

Results

Staphylococcus aureus was observed in 52 (21.1%) samples of 246 raw milk samples. According to the results of PCR assay by amplification of the 23S rRNA gene specific to *S. aureus*, all 52 isolates contained 1250

Table 1: Primers used for the detection of *Staphylococcus aureus* SE genes

410

bp DNA fragments bands and showed positive PCR assay. Table 2 shows the results of molecular tests for the detection of gene encoding the enterotoxins SEA, SEB, SEC, SED, SEE, SEG, SEH, SEI and SEJ. Of the 52 isolates of S. aureus tested, 42 (80.7%) were positive for one or more SE genes, and 12 different genotypes were observed. Among the 52 S. aureus isolates, 28 isolates (53.8%) harbored only one enterotoxin gene, 8 isolates (15.38%) carried gene coding for two enterotoxin. Genotypes encoding three enterotoxins (Sea+Seb+Seh+, Sea+Seb+Sej+, Seb+Sed+Seg+) were detected in 6 isolates (11.53%). Genes encoding the enterotoxins SEC and SEE were not observed separately. Among the genes that code for classic enterotoxins (SEA-SEE), Sea was the most frequent, it was found in 16 isolates (30.7%) followed by Seb in 14 (26.9%) and Sed in 8 (15.37%) isolates. Regarding the other enterotoxins, Seg and Seh were the most frequently observed (8 isolates each, 15.38%), followed by Sej in 6 (11.5%) isolates and Sei in 1 (3.84%) isolate (Fig. 1).

 Table 2: Genotypic profile of Staphylococcus aureus strains isolated from raw milk according to SE genes

Genotypic profile	Number of strains	
Sea	10	
Seb	4	
Sed	4	
Seg	4	
Seh	4	
Sei	2	
Sea+Sed	2	
Seb+Sej	4	
Seg+Seh	2	
Sea+Seb+Seh	2	
Sea+Seb+Sej	2	
Seb+Sed+Seg	2	

Gene	Primer	Sequence (5'-3')	Size (bp) of PCR product	Reference
23S rRNA	23S rRNA F 23S rRNA R	ACG GAG TTA CAA AGG ACG AC AGC TCA GCC TTA ACG AGT AC	1250	Straub <i>et al.</i> 1999
Sea	sea F sea R	TAA GGA GGT GGT GCC TAT GG CAT CGA AAC CAG CCA AAG TT	180	Cremonesi et al. 2005
Seb	Seb F Seb R	TCG CAT CAA ACT GAC AAA CG GCA GGT ACT CTA TAA GTG CC	478	Johnson et al. 1991
Sec	sec F sec R	ACC AGA CCC TAT GCC AGA TG TCC CAT TAT CAA AGT GGT TTC C	371	Cremonesi et al. 2005
Sed	sed F sed R	TCA ATT CAA AAG AAA TGG CTC A TTT TTC CGC GCT GTA TTT TT	339	Cremonesi et al. 2005
See	see F See R	AGG TTT TTT CAC AGG TCA TCC CTT TTT TTT CTT CGG TCA ATC	209	Mehrotra et al. 2000
Seg	seg F seg R	CCA CCT GTT GAA GGA AGA G TGC AGA ACC ATC AAA CTC GT	432	Cremonesi et al. 2005
Seh	seh F seh R	TCA CAT CAT ATG CGA AAG CAG TCG GAC AAT ATT TTT CTG ATC TTT	463	Cremonesi et al. 2005
Sei	sei F sei R	GGT GAT ATT GGT GTA GGT AAC ATC CAT ATT CTT TGC CTT TAC CAG	454	Omoe et al. 2002
Sej	sej F sej R	GGT TTT CAA TGT TCT GGT GGT AAC CAA CGG TTC TTT TGA GG	306	Cremonesi et al. 2005

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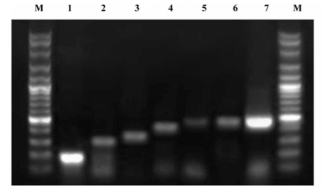


Fig. 1: PCR amplification for the detection of *staphylococcus aureus* SE genes. Lane M: 100 bp marker. Lane 1: sea (180 bp), Lane 2: sej (306 bp), Lane 3: sed (339 bp), Lane 4: seg (432 bp), Lane 5: sei (454 bp), Lane 6: seh (463 bp), and Lane 7: seb (478 bp)

Discussion

Staphylococcus aureus produces a spectrum of extracellular protein toxins and virulence factors which are thought to contribute to the pathogenicity of the organism. The SEs are recognized agents of the Staphylococcal food poisoning syndrome (Straub et al., 1999). In the present work, nine major types of enterotoxin genes are investigated. Considering the genes encoding enterotoxins, 28 (53.8%) out of 52 isolates of S. aureus were positive for at least one enterotoxin gene. The most frequently observed gene was Sea, observed in 16 (30.7%) isolates. In spite of the great discrepancy in data concerning the prevalence of enterotoxigenic S. aureus isolates found in the literature, which is attributable to the different types of foods and strains involved (Mathieu et al., 1991), SEA is the most frequently observed enterotoxin in enterotoxigenic strains of S. aureus (Normanno et al., 2005). In the current study, it was revealed that genes of Sea and Seb, the newly described enterotoxin genes of Seg, Seh and Sej seemed to be the predominant enterotoxin genes of S. aureus isolated from milk of cows. These data were accordance with the finding of Akineden et al. (2001), too. Our study showed that 80.7% of 52 isolates were positive for the presence of genes coding for one or more enterotoxins. In Italy, Morandi et al. (2007) found that 67% of the S. aureus isolated from milk and dairy products were positive for the presence of toxin genes. In Japan, Omoe et al. (2002) observed that 77.4% of the S. aureus isolates were positive for the presence of genes that encode one or more enterotoxins, a frequency close to that observed in the present work (80.7%). With the discovery of the new enterotoxins, the percentage of enterotoxigenic or potentially enterotoxigenic S. aureus isolates increased. In this study, 28 (53.8%) isolates were positive for at least one Se gene, however, that number would decrease to 18 (34.6%) isolates if only the classic enterotoxins (Sea to See) were considered. Rosec and Gigaud (2002) also observed the increase in the number of enterotoxigenic isolates as a consequence of the discovery of the new SEs. In their study, 30% of the

isolates had genes encoding the classic toxins in which frequency was found to be 57% when the new SEs were taken into account. In the present work, six isolates (11.53%) were positive for genes encoding three enterotoxins. Their individual genotypes were *Sea+Seb+Seh+*, *Sea+Seb+Sej* and *Seb+Sed+Seg+*. Nashev *et al.* (2002) have identified genetic profiles comprising multiple genes in *S. aureus*.

In conclusion, the genotypic results of the present study might help to understand the distribution of enterotoxigenic *S. aureus* clones among bovine isolates. This can aid in the investigation and control of *S. aureus* infections in dairy herds.

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مقاله کوتاه: پروفایل های ژنی انترو توکسین در میان Staphylococcus aureus جدا شده از شیر خام

راضیه نظری^۱، حسین گودرزی^۱، فاطمه رحیمی باقی^۱ و مصطفی معینراد^۲

^۱دانشگاه آزاد اسلامی، واحد قم، گروه میکروبیولوژی، دانشکده علوم پایه، قم، ایران؛ ^۲دانشگاه آزاد اسلامی، واحد زنجان، گروه میکروبیولوژی، دانشکده علوم پایـه، زنجان، ایران

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شیر به دلیل دارا بودن مواد مغذی مهم مآنند پرَوتنینها و ویتامینها به عنوان یک غذای مفید در نظر گرفته میشود. همچنین شیر قادر به انتقال باکتریهای پاتوژن مختلف مانند استافیلوکوکوس اورئوس میباشد. در این مطالعه فراوانی ژنهای کد کننده ۹ انتروتوکسین استافیلوکوکی و پروفایلهای ژنی انتروتوکسین در استافیلوکوکوس اورئوس هیباشد. در این مطالعه فراوانی ژنهای کد کننده ۹ انتروتوکسین استافیلوکوکی و پروفایلهای ژنی انتروتوکسین در استافیلوکوکوس اورئوس های جدا شده از شیر خام گاو مورد بررسی قرار گرفت. در مجموع ۲۴۶ نمون به شیر از بر از هشت دامداری مختلف در شهر قم جمعآوری و ۵۲ ایزوله استافیلوکوکوس اورئوس به دست آمد. بر اساس ویژگیهای کشتی، بیوشیمیایی و تکثیر ژن هشت دامداری مختلف در شهر قم جمعآوری و ۵۲ ایزوله استافیلوکوکوس اورئوس به دست آمد. بر اساس ویژگیهای کشتی، بیوشیمیایی و تکثیر ژن همت دامداری مختلف در شهر قم جمعآوری و ۵۲ ایزوله استافیلوکوکوس اورئوس به دست آمد. بر اساس ویژگیهای کشتی، بیوشیمیایی و تکثیر ژن مداله در اساس ویژگیهای کستی، سراند. از نوبی مناسایی شدند. از بیوشیمیایی و تکثیر ژن ۲۰۸۸ ایزولهها، از نظر یک یا چند ژن کد کننده انتروتوکسین مثبت بوده و ۱۲ ژنوتیپ مختلف شناسایی گردید. شایع ترین ژن، ژن کد کننده انتروتوکسین ها به عنوان استافیلوکوکوس اورئوس شاسایی گردید. انیع ترین ژن، ژن کد کننده انتروتوکسینها، شایع ترین زنهای کو که و مور (۸ ایزوله، ۲۰/۹۱)، بود. ژنهای عود و ۱۷ ایزوله، ۲۰/۹۱)، بود. ژنهای Sec در ژن، ژن که در ایلی گردید. ایع ترین ژن، ژنه ای Sec در آنای ژنهای کود در شای ژنهای کود مور در ۱۰ ایزوله، ۲۰/۹۱)، و مور در ۱۱ ایزوله، ۲۰/۹۱)، بود. زنهای عور در ۱۵ (۸ ایزوله، ۲۰/۱۸)، بود. ژنهای عور در مور در در این زنهای که در میر زن که در میر زنده مورد را در میزین ژن، Sec در میلی کردنده مور در ۱۸ (مازله، ۲۰/۱۸)، بود. مورو در مور در ۲۵ (۸ اینوتوکسینهای شایع ترین ژن، Sec در میزولی سویه مولد و مول در هیوکدام از ایزوله، ۲۱/۱۸)، و در مین (۲۰/۱۸ (۱۰ زوله، ۲۰/۱۸)) و در ۱۵ (۸ ایزوله، ۲۰/۱۸)) و مور در این مولد در می ژنهای کور در این در میر در در این مولد در میوتوکسین مولد در میوتوکسین مولد در میوتوکسین مولد در مین مولد در مین مولد در می از آن مور در این موله در مروله، ۲۰/۱۸) و بول در دریل مول در مول در می مول در مول مولوکوکوا مولوکوکوا مول مولی

واژههای کلیدی: Staphylococcus aureus، انتروتوکسینهای استافیلوکوکی، شیر خام، واکنش زنجیرهای پلیمراز