ABSTRACT

Background: *Staphylococcus aureus* is a gram-positive bacterium that has remained a persistent pathogen, causing infections such as endocarditis, meningitis, and toxic shock syndrome in humans. The accessory gene regulator (agr) system of *Staphylococcus aureus* is responsible for controlling the expression of many genes that code for virulence factors. In this study, we assessed the *S. aureus* agr Group, based on their source of isolation, in Gorgan, North of Iran.

Materials and Methods: DNA of 194 *S. aureus* isolates was extracted by lysozyme-phenol chloroform method, which included 85 clinical samples, 58 samples which were isolated from noses of health care workers and 51 cases which were obtained from food products in Gorgan, northern Iran. PCR-based assays were used to evaluate agr locus nucleotide polymorphism for the identification of agr specificity Group. Distributions of each agr Group were determined and comparison between different sources was assessed by X². A p-value of <0.05 was considered as significant.

Results: The majority of isolates belonged to agr Group I (43.3%), followed by agr Group III (28.87%), agr Group II (22.68%), and agr Group IV (5.15%). In our study, a majority of *S. aureus* isolates were recovered from health care workers and food product specimens were of agr Group I and isolates which were recovered from patients were of agr Group III. These differences were statistically significant (P=0.005). There was no statistical difference between the source of isolation of clinical samples of *S. aureus* and agr type.

Conclusion: Agr Group I was predominant among health care workers and food product specimens in Gorgan, North of Iran, but in strains which were isolated from patients, agr Group III was predominant. Investigating the possible role of agr Group III in *Staphylococcus aureus* infection in future studies is recommended.

Keywords: *S. aureus*, Agr Group genes, PCR

INTRODUCTION

*Staphylococcus aureus* is a major cause of both community and hospital-acquired infections; it is a member of the human microbial flora which is responsible for infections which range from subcutaneous abscesses or furuncles, scalded skin syndrome, sepsis, necrotizing pneumonia, pyogenic arthritis and Toxic Shock Syndrome (TSS) [1]. Morbidity/mortality which results from *S. aureus* infection widely varies, depending on the clinical entity, with an incidence rate which ranges from 20 to 50 cases/100,000 population per year, which leads to 10% and 30% deaths [2]. To cause so many human diseases, the accessory gene regulator (agr) globally controls the coordinated production of virulence factors. This system is based on a two-component module which is known as the agr-locus (accessory-gene-regulator), that, in a cell density-dependent manner and through a secreted auto-Inducing-Peptide (AIP), allows a bacterial population to respond in concert when a critical cell-density is reached. The agr locus is composed of two divergent transcriptional units, RNAII and RNAIII, which are driven by the P2 and P3 promoters [3,4].

The P2 operon encodes four proteins (agrA, agrB, agrC, and agrD) and P3 promoter in the opposite direction, encoding the agr system effector molecule (RNAIII). agrD encodes a cyclic AIP that is processed and secreted into the extracellular space via the gene product of agrB. When a critical bacterial cell density is reached, concentrations of AIP bind to the receptor histidine kinase, AgrC, resulting in its activation and subsequent phosphorylation of AgrA. Phosphorylated AgrA then activates transcription of RNAIII at the P3 promoter. RNAIII serves as a transcription factor, turning on expression of genes which encode secreted virulence factors and down regulating expression of cell-associated virulence factors. An AIP with a thiolactonic ring structure, in the early exponential phase, causes immediate activation of the two promoters [4-6].

*Bacterial isolates*: One hundred ninety four isolates of *Staphylococcus aureus* were studied in Golestan University of Medical Sciences. Study samples were collected from health care workers (58 samples), patients (85 samples) and food products (51 samples) between 2009 and 2012, at Gorgan which is located in the north of Iran. The isolates were identified by their growths on Manitol Salt Agar media, Gram staining, catalase test, slide or tube coagulase test, Dnase test and the presence of glutamate synthetase gene [11].

Genomic DNA Extraction: Bacterial DNA lysates were prepared as follows; 1-ml overnight culture of each *S. aureus* isolate was lysed by using lysozyme-phenol chloroform method and it was treated

S. *aureus* isolates can be divided into four agr groups on the basis of the agrC gene which encodes the receptor of the auto inducing peptide and agrD gene, which are responsible for encoding cyclic AIP [7].

Several studies have shown that there was a link between type of agr and the *Staphylococcal* disease. Jarraud and colleagues [8], showed that *Staphylococcus aureus* TSST-1-producing isolates belonged to agr specificity Group III and that most of the exfoliatin-producing strains which were responsible for SSSS belonged to agr Group IV. Ben Ayed and colleagues [9], showed that agr Group III strains were associated with non invasive infections and that agr Group I strains were associated with invasive infections, especially bacteraemia. Chini and colleagues [10] found that TSS toxin 1-producing isolates belonged to agr specificity Groups I and III.

In this study, we investigated the prevalence of agr Groups in *S. aureus* isolates obtained from patients, health care workers and food products, to detect predominant type of isolate, according the source of *S. aureus* and to assess the possible relationship between agr Groups and infection types.
with N-lauroyl sarcosine sodium salt -2% (300 µL), Proteinase K -100 µg[30 µL], and RNase A- 5 µL. DNA was extracted by phenol chloroform, isoamylalcohol, chloroform, and cold ethanol methods.

**Agr typing:** The agr specificity Groups were determined by PCR by using specific primers, which has been shown in [Table/Fig-1] [7].

<table>
<thead>
<tr>
<th>Gene</th>
<th>Primers</th>
<th>Product size</th>
</tr>
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<tbody>
<tr>
<td>agr I</td>
<td>PanF 5-ATG CAC ATG GTG CAC ATG C-3</td>
<td>441-bp</td>
</tr>
<tr>
<td></td>
<td>R 5-GTC ACA AGT ACT ATA AGC TGC GAT-3</td>
<td></td>
</tr>
<tr>
<td>agr II</td>
<td>R 5-TAT TAA TTA TTA AAG GTG GG CAT AGC-3</td>
<td>575-bp</td>
</tr>
<tr>
<td>agr III</td>
<td>R 5-GTA ATG TAA TAG CTT GTA TAA TAA TAC CCA G-3</td>
<td>323-bp</td>
</tr>
<tr>
<td>agr IV</td>
<td>R 5-CGA TTA TGG CGT CGT AAT ACC CG-3</td>
<td>659-bp</td>
</tr>
</tbody>
</table>

![Table/Fig-1](https://www.jcdr.net): The genes and related primers used in this study.

The PCR assay was performed in 25 µL of reaction mixture which contained: 1.5 U of Taq DNA polymerase, 200 µM of dNTPs, 5 mM of MgCl2, 2.5 µL of 10 x PCR buffer, 5 µL of the purified nucleic acid solutions and a 1 µM concentration of each primer. The thermal profile involved an initial denaturation step which was carried out at 94°C for 6 min, followed by 32 cycles of denaturation at 95°C for 45 s, primer annealing at 56°C for 1 min, and primer elongation at 72°C for 70 s. The cycling was followed by a final extension step which was carried out at 72°C for 8 min [7]. PCR products were electrophoresed in a 1.5% agarose gel and they were stained with ethidium bromide. Statistical analyses were done by using SPSS software, version 16 and X2 Statistical test.

**RESULTS**

One hundred ninety four *S. aureus* isolates which were obtained from health care workers, patients and food products, were studied. A majority of isolates belonged to agr Group I (43.3%), followed by agr Group II (22.7%), agr Group III (28.87%), and agr Group IV (5.15%) [Table/Fig-2].

**DISCUSSION**

*Staphylococcus aureus* is a major cause of both community and hospital acquired infections; it is a member of the human microbial flora which is responsible for infections which range from subcutaneous abscesses or furuncles, to scalded skin syndrome, sepsis necrotizing pneumonia, and toxic shock syndrome (TSS). Many cell surface proteins which are secreted, exotoxins, enzymes and virulence factors of *S. aureus*, are regulated by agr locus [1].

*S. aureus* isolates can be divided into four agr Groups on the basis of the agrC gene which encodes the receptor of the auto inducing peptide and agrD gene which encodes the auto inducing peptide [7]. In our study, *Staphylococcus aureus* was classified, based on agr locus in four agr Groups. Dufour and colleagues [12], first used this method for classification of *Staphylococcus aureus* and they showed that isolates of this bacterium could be divided into four Groups, I, II, III, IV. Although agr specific Group IV was absent in many previously reported studies [7,13,14], we detected agr Group IV in blood, wound and urine samples. As was seen in a majority of previous studies, in our region too, agr Group I was the most prevalent agr type. For example, Shopsin and colleagues [7] found that agr specific Group I (42%) was prevalent in children and their guardians. In the study done by van Leeuwen and colleagues’ [13] on a collection of 192 *S. aureus* strains, 71% of strains were found to belong to agr Group I and in the study done by Najar and colleagues [15] on a collection of 212 *S. aureus* strains, 55.1% of strains were found to belong to agr Group I. In a more recent study done by Indrawattana and colleagues in 2013 in Thailand, it was found that agr specific Group I (58.7%) was predominant agr Group [16].

The predominant agr type which was isolated from food products in present study was agr I, followed by agr Group II, but in a study which was conducted by Montaz in 2010, agr Group II was found to be most prevalent among *S. aureus* which was isolated from milk in Iran [17].

The predominant agr Group found in patients was Group III, but it was less frequent in healthcare workers and food products. Ben and colleagues, showed in Tunis [9] that among a total of 57 *S. aureus* strains which were isolated from patients, 9 (15.7%) belonged to Group I, 2 (3.5%) belonged to Group II and that 23