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Original Research Article

Virulence factors of Methicillin Resistant Staphylococcus aureus (MRSA) isolated from burn patients

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ABSTRACT

	The present study aimed to investigate some of virulence factors among Methicillin Resistant <i>Stanbylococcus aureus</i> (MRSA) isolated from hurn wound. From a total					
Keywords	of 126 isolates of <i>S. aureus</i> , only eighty five(67.46%) isolates of MRSA were					
MRSA, Burn	July to November, 2014 in Thi-Qar province, Iraq.All MRSA isolates were examined using Polymerase Chain Reaction (PCR) for detection 16SrRNA mec.					
Virulence	gene and some virulence factors of this bacteria include <i>sea</i> , <i>hla</i> , <i>hlb</i> , and <i>cap</i> 8 The results revealed that all isolates have 16SrRNA and <i>mec</i> A genes that were					
Molecular detection	used to confirm these bacteria as Staphylococci and MRSA respectively. The virulence factors detection results showed percentages of (72.941%), (82.352%)					
	and (85.882 %) of isolates have <i>sea</i> , <i>hla</i> and <i>hlb</i> genes respectively, while only 69 (81.176%) of isolates have <i>cap</i> 8 gene.					

Introduction

Thermal injury destroys the skin barriers that normally prevent invasion by microorganisms (Singh *et al.*, 2003). Burn patients become susceptible to various infections due to the loss of this protective barrier and decreased cellular and humoral immunity (Wong *et al.*, 2002). In these patients, burn wound infections can easily escalate into sepsis (Church *et al.*,2006).

The common pathogens isolated from burn wound are *S.aureus*(75%), *Pseudomonas aeruginosa*(25%), *Streptococcus pyogenes* (20%) and various coliform bacilli (5%) (Ahmad and Iranzo, 2003).

MRSA is the most important pathogen among Staphylococi (Lee et al., 2007). MRSA strains are isolated in more than half of all community and hospital infections (Klevens et al., 2007). MRSA has become a major public health problem worldwide, and the problem of MRSA continues to rise (Nimmo et al., 2006 and Jarvis et al., 2007). MRSA has been the most commonly recognized multidrug-resistant pathogen in the universe and the emergence of MRSA strains found in increasing number of infections and often multi drug resistant in nature now pose serious therapeutic problems to clinicians (Groundmann et

al.,2006; Marais *et al.*,2009).

Most MRSA strains carry *mec*A encoding low affinity penicillin-binding protein PBP2a (or PBP2') (Hiramatsu *et al.*, 2001). The MRSA characteristic phenotype is due to the presence of *mec* Awhich encodes a PBP2a, with degraded affinity for β -lactams (Oliveira and De Lencastre, 2011; Moellering, 2012).

The virulence factors of *Staphylococcus* include surface components, such as the peptidoglycans, teichoic capsule, acid. protein A, enzymes such as (esterases, lipases, fatty-acid modifying enzymes, hydrolytic various proteases, enzymes, catalase, betalactamase), and various toxins, such as (leukocidins, enterotoxins, TSST-1 and alpha, beta, gamma and delta (Vasconcelos and Cunha, hemolysins) 2010).

One of the virulence factors of S. aureus is cytolytic, pore-forming toxin (Diep and Otto, 2008), such as alphahemolysin (Hla) has been implicated in the pathogenesis of S. aureus (Labandeira-Rey et al., 2007). Hla has cytolytic activity toward a variety of host cell types, including human epithelial keratinocytes. cells and lymphocytes (Hocke al.. 2006; et Wardenburget al., 2008). Other types of hemolysin is Beta-hemolysin (Hlb) is a magnesium-dependent sphingomyelinase C that induces lysis of sheep erythrocytes and human monocytes (Walev et al., 1996).

S. aureus is produced one of the extracellular protein toxins, staphylococcal heat stable enterotoxin (SE) is the most important virulent factors belonging to the superantigen family (Pinchuk *et al*, 2010), and many strain of *S. aureus*, especially MRSA, secreted one or more specific staphylococcal exotoxins, including

staphylococcal enterotoxins (SEs), (Llewelyn and Cohen, 2002).

Among those factors considered for typing, capsular polysaccharides expressed by *S. aureus* are one of them, since they are also important in the pathogenesis of staphylococcal infections, most *S. aureus* isolates are encapsulated and so far eleven capsular serotypes have been described, of these, types *cap* 5 and *cap* 8 predominate in approximately 75% of the clinical isolates (Murphy *et al.*, 2011).

Materials and Methods

Samples collection

Two hundred and seventy six samples were collected from burn patients in burn unit of AL-Hussain Teaching Hospital of Thi-Qar province in the period from July to November, 2014 by moistened sterile swabs with normal saline, then these swabs directly inoculated on Mannitol salt agar (LAB/United Kingdom) and incubated at 37°C for 24 hours.

Identification of *S. aureus*

S. aureus was identified depending on the morphological properties on culture media and biochemical tests which done according to Bergeys manual (MacFaddin, 2000).API Staph system was used as identification system for *Staphylococcus* and *Micrococcus*. This test was done according to the company instructions (BioMerieux, France).

StaphyloMonotec test kit Plus

This kit is a new rapid agglutination test for differentiation between *S. aureus* and other *Staphylococcus*. The test was performed according to the directions of manufacturing

company (Fluka Analytical, Switzerland).

Antibiotic sensitivity test

The antibiotic sensitivity test was done by the agar disc diffusion method as described by (Kirby and Bauer, 1966).

Molecular Detection

DNA extraction

DNA from all MRSA isolates were extracted using Genomic DNA Extractionspin kit (Bosphore, Anatolia genewors). 16S rRNA, *mecA*, *sea*, *hla*, *hlb*, and *cap*8 genes were identified by using primers describedin Table (1).

Amplification of the mecA gene was done using primer described by (Jonas et al., 2002). The final volume of reaction tubes is 20µl, consist of 10 µl Master Mix., 1.25µl of both Forward (F) and Reverse (R) of the primer specific for the mecA gene, 5µl of template DNA and complete the volume by adding free water to 20µl.Amplification of the 16SrRNA, sea, hla, hlb, hla, and cap8 genes were done using primers described as above (Table1). The final volume of reaction tubes is 20µl, consist of 10 µl Master Mix., 1µl of both F. and R. of the primers specific for these genes, 5µl of template DNA and complete the volume by adding deionizing water to 20µl.

Results and Discussion

Eighty five isolates (67.460 %) of MRSA were obtained from *S. aureus* isolates that collected from burn patients. All strains identified was done by cultural, biochemical and serological tests to confirm exact identification as*S. aureus*. MRSA outbreaks are estimated in about 40-60% of *S. aureus* outbreaks, which are mainly affected by the

infection control program and medical treatments leading to a wide range of hospital infections (Fatholahzadeh *et al.*,2008). Alfatemi *et al.*,(2014) showed the prevalence of MRSA among *S. aureus* isolates was (42.3%), which indicates little difference in terms of frequency with studies by Fatholahzadeh *et al.*,(2008) who reported MRSA prevalence of (36%) in Tehran.

The prevalence of MRSA in present study wasslightely, in agreement with other studies in Iraq, which recorded percentages of (65.3%), (88%) and (75%) respectively (Al-Mussawi, 2014; Yaseen *et al.*,2013 and Al-Azawi, 2013).

The molecular diagnostic of this bacteria was performing to all MRSA isolates through the amplification of 16SrRNA and *mec* A genes to confirm that the tested isolates are staphylococci and MRSA respectively. Other genes were used to detect many virulence factors of MRSA that included *hla*, *hlb*, *sea* and *cap*8 genes.

All isolates were showed positive results of and *mec* A genes both 16SrRNA (100%)(Table 2). The bands were 756, 310 bp size corresponds to amplification of16SrRNA and mecA genes respectively, Fig(1 and 2).Al-Talib et al.,(2009)reported that all isolates had 16SrRNA, 82 contained mecA genes. Current study results agreed with Makgotlho, (2009) who showed that all isolates 97/97 (100%) have 16SrRNA gene while mec A gene was detected in 96/97 (99%) of the MRSA isolates, which did not show the presence of mec A gene was, however phenotypically identified as MRSA.

The results showed that 69 (81.1 %) of isolates have cap8 gene (Table 2), the bands were (450 bp) size corresponds to amplification of cap8 gene, Fig (6). Udo and

Sarkhoo (2010) reported that capsular polysaccharides and types prevalence was (77.3%) and only three isolates (2.2%) yielded negative result for both*cap5* and *cap8*.

Seventy (82.35%) of isolates have *hla* gene only, Table (2). The bands were (209 bp) size corresponds to amplification of hla gene, Fig (4). Most of S. aureus isolated from human have usually an alpha haemolytic character, because the human platelets and monocytes are more sensitive to the alpha toxin (Todar, 2005). Kateete et al.(2011) showed the frequency of hlagene was 100%. Likewise, in a study from the United States the *hlagene* frequency was reported at 100% (Shukla et al., 2010). The percentage of *hlb* gene in MRSA isolateswas 73 (85.88%), Table (2). The bands were

(833 bp) size corresponds to amplification of hlb gene, Fig (5). The study performed by Rusenova et al.,(2013) showed that 31 MRSA isolates (42.5%) for beta toxin, 41 (56.2%)of isolates showed double hemolysis (alpha + beta hemolysins), and 1 (1.4%) was non-hemolytic. MRSA isolates have 62 (72.94%) of sea gene Table (2). The bands were (120 bp)size corresponds to amplification of sea gene, Fig (3). Alfatemi et al.,(2014) the frequency of the sea gene was 27.39%. Our study was in agreement Sarkhoo, with Udo and (2010)that 103 (76.3%) isolates whomreported yielded positive results for sea. However, the role of S. aureus superantigenic toxins in the severity of septicemia patients should not be discounted as sea is significantly associated with severity of sepsis caused by S. aureus (Ferry et al, 2005).

Table.1 Oligonucleotide primers sequences for PCR amplified of 16SrRNA,mecA, sea, hla, hlb and cap 8 genes

Genename	Primer Sequences (5'- 3')	Length	References
	F: AAC TCT GTT ATT AGG GAA GAA CA		
16SrRNA		756 bp	(McClure et al., 2006)
	R: CCA CCT TCC TCC GGT TTG TCA CC		
	F: GTA GAA ATG ACT GAA CGT CCG ATA A	2101	(C. 1 1. 1004)
mecA		310 bp	(Gena <i>et al.</i> , 1994)
hlb	F' GCC AAA GCC GAA TCT AAG		
		833 bp	(Booth et al., 2001)
	R: GCG ATA TAC ATC CCA TGG C	1	· · · /
hla	F: CTG ATT ACT ATC CAA GAA ATT CGA TTG		
		209 bp	(Mehrotraet al.,2000)
	R: CTT TCC AGC CTA CTT TTT TAT CAG T		
cap8	F: GCG CTA CAA ACA TTA AGC AT	1501	(0 1 1007)
		450 bp	(Sau <i>et al.</i> ,1997)
	R: TIC TIA GCC IGC IGG CAT C		
			(Datlay and Makalana-
sea		120 bp	(Belley and Mekalanos,
	R: GAACUTTCCCATCAAAAACA		1988) with modified

Int.J.Curr.Microbiol.App.Sci (2015) 4(7): 898-906

Genes	Positive %	Negative %
16SrRNA	85 (100 %)	-
mecA	85 (100 %)	-
hla	70 (82.352 %)	15 (17.647 %)
hlb	73 (85.882 %)	12 (14.117 %)
sea	62 (72.941 %)	23 (27.058 %)
cap 8	69 (81.176 %)	16 (18.823 %)

Table.2 The percentage of genes in MRSA isolates





16srRNA (756bp)

Figure.2 agarose gel electrophoresis of mec A gene amplification





Figure.3 Agarose gel electrophoresis of sea gene amplification

Figure.4 Agarose gel electrophoresis of hla gene amplification



Figure.5 Agarose gel electrophoresis of hlb hemolysin gene amplification





Figure.6 Agarose gel electrophoresis of cap8 gene amplification

References

- Ahmad, S.I and Iranzo, O. G. (2003). Treatment of post burns bacterial infections by Fenton reagent, particularly the ubiquitous multipledrug resistant *Pseudomonas* species. *Medical Hypotheses*, 61(4):431–434.
- Al-Azawi, I.H.S. (2013). Antibiotic susceptibility pattern and *mec* A gene detection in methicillin resistant *Staphylococcus aureus* (MRSA) isolated from burn and wound in Al-Diwaniya city. *Journal of Babylon university*, Pure and applied sciences; 21(3): 917-926.
- Al-Mussawi, A.A. (2014). Detection of Staphylococcus aureus and Methicillin Resistant Staphylococcus aureus (MRSA) from human clinical specimens using conventional biochemical tests and chromogenic media. Indian Journal of applied research ;4 (2):7-9.
- Al-Talib, A.; Yean, C.Y.; Al-Khateeb, A.; Hassan, H.; Singh, K.B.; Al-Jashamy, K. and Ravichandran, M. (2009). A pentaplex PCR assay for the rapid detection of Methicillin-Resistant *Staphylococcus aureus*and Panton-Valentine Leucocidin. *BMC Microbiology*, 9:113 doi:10.1186/1471-2180-9-113.
- Betley, M.J. and Mekalanos, J.J. (1988). Nucleotide sequence of the typeA

staphylococcal enterotoxin gene. Journal of Bacteriology 170, 34–41.

- Booth, M.C.; Pence, L.M.; Mahasresthi,P.; Callegan, M.C. and Gilmore, M.S. (2001). Clonal association among *Staphylococcus aureus* isolates from various sites of infection. *Inf. and Immun*, 69: 345-352.
- Church, D.; Elsayed, S.; Reid, O.; Winston, B. and Lindsay, R. (2006). Burn woundinfections. *ClinMicrobiol Rev*, 19: 403–434.
- Diep, BA. and Otto, M. (2008). The role of virulence determinants in community associated MRSA pathogenesis. *Trends Microbiol*, 16: 361–369.
- Fatholahzadeh, B.; Emaneini, M.; Gilbert, E.;Aligholi, M.;Modarressi, G.;Udo, MH.et al. (2008).Staphylococcal cassette chromosome mec (SCCmec) analysis and antimicrobial susceptibility Resistant patterns of Methicillin Staphylococcus aureus (MRSA) isolates in Tehran, Iran. Microb Drug Resist.;14(3):217-20.
- Ferry, T.; Thomas, D.;Genestier, AL., *et al.* (2005). Comparative prevalence of superantigen genes in *Staphylococcus aureus* isolates causing sepsis with and without septic shock. *Clin Infect Dis*, 41: 771-7.
- Geha, D. J.; Uhl,J. R.;Gustaferro,C. A. and Persing, D. H. (1994). Multiplex PCR

for identification of Methicillin Resistant staphylococci in the clinical laboratory. *J. Clin. Microbiol.* 32:1768–1772.

- Groundmann, H.; Sousa, M.; Boyce, J. and Tiemesma, E. (2006).Emergence and resurgences of MRSA as a public health threat. *Lancet.*; 368(9538):874-85.
 Hiramatsu, K.; Cui, L.; Kuroda, M. and Ito, T. (2001). The emergence and evolution of Methicillin Resistant *Staphylococcus aureus. Trends Microbiol*; 9: 486-93.
- Hocke, AC.; Temmesfeld-Wollbrueck, B.: Schmeck, B.et al. (2006). Perturbation of endothelial junction proteins bv Staphylococcus aureus alpha-toxin: inhibition of endothelial gap formation by adrenomedullin. J Histochem Cell 126:305-316. Biol: Alfatemi, H.: S.M.H.; Motamedifar, M.; Hadi, N. and Saraie, H.S.E. (2014). Analysis of Virulence Genes Among Methicillin *Staphylococcus* Resistant aureus (MRSA) Strains. JundishapurJ Microbiol. 7(6): e10741.
- WR.: Schlosser. J.: Chinn, Jarvis. RY.;Tweeten, S. and Jackson, M. (2007).National prevalence of methicillin-resistant *Staphylococcus* aureus in inpatients at US health care facilities, Am. J. Infect. Control., 35: 631-637.
- Jonas, D.; Speck,M.; Daschner, F. D. and Grundmann, H.(2002). Rapid PCR-Based Identification of Methicillin-Resistant *Staphylococcus aureus* from Screening Swabs. J. Clin. Microbiol., 40(5):1821-1823.
- Kateete, DP.; Namazzi, S.; Okee, M.; Okeng, A.;Baluku, H.; Musisi, NL.et al. (2011).
 High prevalence of methicillin resistant Staphylococcus aureus in the surgical units of Mulago hospital in Kampala, Uganda. BMC Res Notes;4:326.
- Kirby, W.M.; Baur, A.W.;Scherris, J.C.;and Torch, M. (1966). Antibiotic susceptibility testing by standardized single methods. AM. *J.eli. Path* 45:493-496.

- Klevens, RM.; Morrison, MA.; Nadle, J. *et al.* (2007). Invasive Methicillin Resistant *Staphylococcus aureus* infections in the United States. *JAMA*, 298: 1763–1771.
- Labandeira-Rey, M.;Couzon, F.;Boisset, S.; Brown, EL.; Bes, M.et al. (2007).*Staphylococcusaureus* Panton-Valentine leukocidin causes necrotizing pneumonia. *J Science* 315: 1130–1133.
- Lee, SS.; Kim, HS.; Kang, HJ.; Kim, JK. and Chung, DR. (2007). Rapid spread of Methicillin Resistant *Staphylococcus aureus* in a new hospital in the broadspectrum antibiotic era. *J Infect*,55: 358-62.
- Llewelyn, M. and Cohen, J. (2002). Superantigens: microbial agents that corrupt immunity. *Lancet Infect Dis* 2, 156–162.
- MacFaddin, J.F.(2000). Biochemical tests for identification of medical bacteria, 3 rd ed.; Lippincott Williams & Wilkins: USA.
- Makgotlho, P. E. (2009). Molecular Characterization of Methicillin Resistant *Staphylococcusaureus* strains, MSC. University of Pretoria, South Africa. Marais, E.;Aithma, N.;Perovic, O.;Oosthuysen, WF.;Musenge, E.;Dusé, AG. (2009). Antimicrobial susceptibility of Methicillin Resistant *Staphylococcus aureus* isolates from South Africa. J S Afr Med, 99(3):170-3.
- McClure, J. *et al.*,(2006). Novel multiplex PCR assay for detection of the Staphylococcal virulence marker pantone valentine leukocidin genes and simultaneous discrimination of methicillin- susceptible from resistant staphylococci. *J of clinic Micro*, 44:1141-4.
- Mehrotra M,; Wang, G. and Johnson, WM. (2000). Multiplex PCR for detection of genes for Staphylococcus aureus enterotoxins, exfoliative toxins, toxic shock syndrome toxin 1, and methicillin resistance. *J Clin Microbiol*.;38(3):1032– 5.

Moellering, Jr. (2012). RC. MRSA: the first

half century. *J Antimicrob.Chemother*, 67:4-11.

- Murphy, E.; Lin, SL.; Nunez,L.; Andrew, L.; Fink, PS.; Dilts, DA.*et al.*,(2011). Challenges for the evaluation of *Staphylococcus aureus* protein based vaccines: monitoring antigenic diversity. *J of Hum Vaccin*; 7:51-9.
- Nimmo, GR.; Coombs, GW.; Pearson, JC.;
 O'Brien, FG. and Christiansen, KJ. (2006). Methicillin Resistant *Staphylococcus aureus* in the Australian community: an evolving epidemic. *Med. J. Aust.* 184: 384-388. 17.
- Oliveira, DC. and De Lencastre, H. (2011). Methicillin-Resistance in *Staphylococcus aureus* is not affected by the Overexpression in Trans of the *mecA* Gene Repressor: A- Surprising Observation. J. PLOS ONE; 6: 1-9.
- Pinchuk IV, Beswick EJ, Reyes VE. (2010). Staphylococcal enterotoxins. *Toxins*, 2:2177-97.
- Rusenova, N.; Gebreyes, W.; Koleva, M.; Mitev, J.; Penev, T. Vasilev, N. and Miteva, T. (2013). Comparison of Three Methods for Routine Detection of *Staphylococcus aureus* Isolated from Bovine Mastitis. J of KafkasUniv Vet FakDerg, 19 (4): 709-712.
- Sau, S.;Bhasin, N. ; Wann,E.R. ; C. Lee, J.C. Foster, T.J. and Lee, C.W. (1997). The *Staphylococcus aureus* allelic genetic loci for serotype 5 and 8 capsule expression contain the type-specific genes flanked by common genes. Microbiology, 143,2395-2405.
- Shukla, SK.; Karow, ME.; Brady, JM.; Stemper, ME.;Kislow, J.; Moore, N.*et al.* (2010). Virulence genes and genotypic associations in nasal carriage, community-associated methicillinsusceptible and methicillin-resistant USA400 *Staphylococcus aureus* isolates. *J Clin Microbiol.*;48(10):3582–92.
- Singh, N.P.;Goyal, R.;Manchanda, V.; Das, S.;Kaur, I. and Talwar, V. (2003). Changing trends in bacteriology of burns in the burns unit, Delhi, India. *Burns*,

29(2): 129-132.

- Todar, K. (2005). Todar Online Textbook of Bacteriology. *Staphylococcus*. University of Wincosin-Madison Department of Bacteriology. www.text bookofbacteriology.net/staph.html.
- Udo, E.E. and Sarkhoo,E. (2010). The dissemination of ST80-SCCmec-IV community-associated methicillin resistant Staphylococcus aureus clone in Kuwait hospitals. Udo and Sarkhoo Annals of Clinical Microbiology and Antimicrobials, 9:31.
- Vasconcelos, NG. and Cunha, MLRS. (2010). Staphylococcal enterotoxins: Molecular aspects and detection methods. *J Public Health Epidemiol*; 2: 29-42.
- Walev, I.; Weller, U.;Strauch, S.; Foster, T. and Bhakdi, S. (1996). Selective killing of human monocytes and cytokine release provoked by sphingomyelinase (beta-toxin) of *Staphylococcus aureus*. *Infect Immun* 64: 2974-2979. Wardenburg, B. J. and Schneewind, O.(2008). Vaccine protection against *Staphylococcus aureus* pneumonia. *J Exp Med*; 205:287–294.
- Wong, T.H.; Tan, B.J.; Ling, M.L. and Song, C. (2002). Multi-resistant Acinetobacter baumannii on a burns unit – clinical risk factors and prognosis. Burns, 28(4): 349-357.
- Yaseen, I.H. *et al.*, (2013). high prevalence of multidrug resistant MRSA and VRSA of different infection from Al-Jumhuory Teaching Hospital patients in Mosul. *J of Life Sciences*, 7(12)1255-9.