Genetic regulation of fatty acid modifying enzyme from *Staphylococcus aureus*

N. R. CHAMBERLAIN and B. IMANOE

*Department of Microbiology/Immunology, Kirksville College of Osteopathic Medicine, Kirksville, MO 63501, USA*

Fatty acid modifying enzyme (FAME) is an extracellular enzyme that inactivates staphylocidal lipids by catalysing the esterification of these lipids to cholesterol. In-vitro expression of FAME began at the start of the stationary phase. This expression of FAME was very similar to other staphylococcal extracellular proteins controlled by the global regulators Agr and Sar. A *Staphylococcus aureus* strain ISP546 (Agr⁻) produced c. 80% less FAME than an isogenic Agr⁺ strain ISP479C. Similar results were obtained with the isogenic Agr⁺/Agr⁻ strain pair RN6390 and RN6911. A *S. aureus* strain R (Sar⁻) produced c. 86% less FAME than an isogenic Sar⁺ strain RN6390. However, lipase assays on the same culture filtrates from the Sar⁺/Sar⁻ strains did not demonstrate any affect on lipase production by the sar mutation.

Introduction

Certain neutral lipids are bactericidal to gram-positive organisms in vitro [1–3]. These bactericidal neutral lipids include long-chain fatty acids and monoglycerides. Previous studies have also demonstrated that the elimination of *Staphylococcus aureus* within murine abscesses is mediated by the production of long-chain fatty acids and 1- and 2-monoglycerides [1–3]. Long-chain fatty acids and monoglycerides comprised c. 40% (by dry weight) of the neutral lipids obtained from an abscess [4]. The amount of neutral lipids found in *S. aureus* murine abscesses [4] (90% neutral lipids; 10% phosphatides and glycolipids) is much higher than is observed in normal tissues [5] (15–40% neutral lipids; 60–85% phosphatides and glycolipids). Production of these neutral lipids occurs c. 24 h after abscess formation [1].

Fatty acid modifying enzyme (FAME) is an extracellular staphylococcal protein that has been shown to inactivate the bactericidal activity of fatty acids [2, 6]. FAME catalyses the esterification of alcohols (e.g., cholesterol, ethanol) with fatty acids [6] (e.g., palmitate, laureate, oleate). The esters resulting from this reaction are not bactericidal. About 80% of the *S. aureus* strains tested thus far produce FAME [7]. It has also been shown that FAME-producing strains of *S. aureus* are more invasive in a murine model [6].

Other investigators have demonstrated the importance of global regulators in controlling the expression of several staphylococcal extracellular proteins [8–20]. Two of these regulatory elements, the accessory gene regulator (Agr) and the staphylococcal accessory regulator (Sar), control expression of exoproteins by transcriptional control [13, 14]. Several of these extracellular proteins are important virulence factors (e.g., α toxin, enterotoxins, toxic shock syndrome toxin) and the timing of expression of these factors may prolong the organism's ability to survive in the host.

Mutations in the Agr locus eliminate expression of some of these exoproteins (e.g., α toxin, serine protease, δ toxin and toxic shock syndrome toxin). These proteins are called class I Agr-regulated proteins. Other exoproteins are still produced in the Agr⁻ strains but at much lower levels (e.g., nuclease, lipase, β toxin and enterotoxin B) and are called class II Agr-regulated proteins. This study examined the effects of mutation of the agr or sar locus on FAME expression and showed that it is a class II Agr-regulated protein.

Materials and methods

Bacterial strains

*S. aureus* strain 18Z was a kind gift from Dr F. A. Kapral and has been characterised previously [21]. *S. aureus* strains ISP479C (Agr⁺), ISP546 (Agr⁻), RN6390 (Agr⁺) and RN6911 (Agr⁻) were used to determine the role of the accessory gene regulator (Agr) in FAME production. The above isogenic strain pairs were kind gifts from Dr J. J. Iandolo (ISP479C/
ISP546) [11, 20] and Dr R. P. Novick (RN6390/ RN6911) [16, 22]. In summary, the Agr− strains produce greatly reduced amounts of lipase, α toxin, protease and nuclease. The Agr− strain (ISP546) was made by insertion of the transposon Tn551 in the agr locus. The Agr+ strain (ISP479C) was obtained by curing ISP546 of the transposon in landolo's laboratory. Removal of the entire agr locus from RN6390 and replacing it with a tetracycline resistance gene resulted in an Agr− strain, RN6911 [16, 22].

S. aureus mutant R Sar− was a kind gift from Dr A. L. Cheung [13, 14]. The mutant R strain was produced by placing Tn917LTV1 in the sar locus of RN6390 (Sar').

In all three strain pairs, α toxin was expressed in the Sar+ and Agr+ strains but was not detected in the Sar− and Agr− strains as has been reported previously (data not shown) [8, 16, 18].

**FAME assay**

The samples (5 μl) to be assayed were diluted with 95 μl of 20 mM MES, 170 mM NaCl, pH 6.0 (MES-NaCl) unless otherwise indicated. To the diluted filtrate (100 μl) was added 5 μl of acetone containing [7-3H]cholesterol (200 000 dpm; sp. act. = 23.8 Ci/ mmol; New England Nuclear, Wilmington, DE, USA) and oleic acid 2.5 μg. Samples were incubated for 30 min at 37°C. The lipids were then extracted from the solution with 200 μl of ethyl ether:methanol (6:1, EE:M). The lower phase was discarded and the upper organic phase was dried in a stream of nitrogen gas. The dried lipids were then suspended in 100 μl of hexane:ethyl ether:glacial acetic acid (73:25:2; H:EE:AA). The cholesterol ester was separated from the radiolabelled cholesterol with silica gel columns and a solvent system used for TLC to separate cholesterol esters from fatty acids and cholesterol as described previously [5, 23]. Slurries of silica gel (average particle size 40 μm; VWR Scientific, St Louis, MO, USA) in H:EE:AA were used to make 0.6-g columns (dry weight; 5.3 cm × 0.5 cm) in 23-cm Pasteur pipettes plugged with siliconised glass wool. The suspended samples were placed on the column and the cholesterol esters were eluted in 2 ml of H:EE:AA. The eluant was collected in liquid scintillation vials and 10 ml of scintillation fluid (ScintiSafe Econo F; Fisher, St Louis, MO, USA) were added. Radioactivity (cpm) of the samples was measured in a liquid scintillation counter as a direct measure of FAME activity (esterification of cholesterol with oleic acid).

**Effects of agr and sar mutations on FAME production**

Strains ISP479C (Agr+), ISP546 (Agr−), RN6390 (Agr+) and RN6911 (Agr−) were used to determine the role of agr in FAME production. The bacteria from an overnight culture (0.5 ml) were pelleted by centrifugation (12 000 g, 10 min, 4°C) and then washed with sterile Trypticase Soy Broth (TSB, Difco) 0.5 ml. This was repeated twice before addition of the organisms to pre-warmed (37°C) TSB (50 ml) in a 250-ml Erlenmeyer flask. The cultures were incubated at 37°C and aerated by shaking at 110 rpm (reciprocating shaker). At various times, 1 ml samples of the cultures were removed, the optical density at 600 nm (OD600) was determined, and filtrate was collected (0.2 μm; Acrodisc; Gelman, Ann Arbor, MI, USA). The filtrates were then stored at −20°C until they were assayed for FAME activity as described above.

Strains RN6390 (Sar+), and R (Sar−) were incubated and assayed for FAME activity as described above. The Sar+ /Sar− strain pair was also assayed for lipase activity as described below. Antibiotics were used only to passage the Sar− and Agr− strains.

**Lipase assay**

The assay was performed as previously described with thin layer chromatography Silica Gel H plates (Analtech; Newark, DE, USA) to separate products from substrate [24].

**Results**

**Genetic regulation of FAME expression**

All three isogenic strain pairs were incubated in TSB at 37°C. Samples were taken at various time intervals to determine when and how much FAME activity was present in the culture filtrates.

The Agr+/Agr− strains ISP546/ISP479C both grew exponentially for 2 h followed by a slower growth phase for about another 4 h. Growth of both strains reached a plateau at c. 6 h. There was no significant difference between the growth curves of the Agr− and Agr+ strains (Fig. 1).

Assays for FAME activity in the culture filtrates were then conducted to see if mutation in the agr locus had any effect on FAME production. FAME activity expressed by the Agr+ strain (ISP479C) increased rapidly from 2 to 7 h of growth (Fig. 1). A plateau was reached at 7–9 h followed by a decrease in FAME activity after growth for 9 h. The greatest increase in FAME activity began during early stationary phase (Fig. 1). Similar results were obtained with a wild-type FAME-producing strain of S. aureus 18Z [21] (data not shown).

The Agr− strain (ISP546) expressed little FAME activity until c. 7 h of growth (Fig. 1). FAME activity did increase between 1 and 2 h of incubation although the amount of FAME produced was less than that produced by the Agr+ strain. Significant increases in
Fig. 1. FAME activity of *S. aureus* strains ISP479C (Agr'; ■) and ISP546 (Agr-; ●) versus time of incubation and the growth curve of ISP479C (Agr'; □) and ISP546 (Agr-; ○). FAME production by the Agr strain did not occur until late in the stationary phase. When FAME activities at 8 h were compared, the Agr- strain expressed 84% less FAME than the Agr' strain.

Similar results were obtained with the other Agr-/ Agr' strain pair (RN6911/RN6390). This Agr' strain also did not express FAME until early in stationary phase (Fig. 2). The Agr- strain began expression of FAME in early stationary phase (2-3 h) although the rate and amount of FAME produced were less. An increase in FAME production by the Agr- strain was observed late in stationary phase. At 8 h of growth the Agr- strain expressed 50% less FAME than the Agr' strain. The reduction in FAME expression that occurred in the stationary phase was not significant.

The Sar-/Sar' strain pair (R/RN6390) was then

Fig. 2. FAME activity of *S. aureus* strains RN6390 (Agr'; ■) and RN6911 (Agr-; ●) versus time of incubation and the growth curve of RN6390 (Agr'; □) and RN6911 (Agr-; ○).
examined to see what affect, if any, this global regulator had on FAME production. The Sar- mutant expressed FAME activity in early stationary phase; however, the rate and amount of FAME produced were less than those produced by the Sar+ strain (Fig. 3). After incubation for 8 h the Sar- strain produced 86% less FAME than the Sar+ strain. Lipase activity after incubation for 8 h was not affected (data not shown).

Discussion

These studies show that FAME expression is significantly reduced in Agr- and Sar- strains, and also demonstrate that FAME is a class II Agr-regulated exoprotein.

Previous studies have demonstrated the importance of certain staphylocidal neutral lipids (fatty acids and monoglycerides) in destroying *S. aureus* in murine abscesses [1–3]. The staphylocidal lipids represent c. 60% (by weight) of the neutral lipid extracted from abscesses. Shortly after abscess formation (24 h) these staphylocidal lipids are produced by the host [3]. One way *S. aureus* may prolong its survival in an abscess is to inactivate the staphylocoidal lipids.

FAME has been shown to protect *S. aureus* against the bactericidal effects of fatty acids by esterification of the staphylocoidal lipids to cholesterol [6]. Previous studies have also shown a positive correlation between the ability of a strain to produce FAME and its ability to cause invasive disease [6].

This study was conducted to determine whether FAME expression was controlled by a staphylococcal global regulator. If FAME is important in the survival of staphylococci in an abscess then the timing of expression may be as important as the amount of FAME produced. As mentioned earlier, staphylocidal fatty acids do not occur in an abscess until about 24 h after abscess formation. To save valuable energy, it could possibly benefit the bacterium to delay FAME production until just before it is needed most.

At least three different global regulators have been described: the accessory gene regulator (Agr) [16–19], staphylococcal accessory gene regulator (Sar) [13, 14] and the extracellular gene regulator (Xpr) [20]. Each one is able to delay the expression of many extracellular proteins until the early stationary phase of growth. The Agr and Sar global regulators have been very well described [13, 14, 16–19]. Isogenic mutants and wild-type strains are readily available and, therefore, were used in this study.

In these experiments it was demonstrated that mutation of the *agr* locus drastically affects FAME production (Figs 1 and 2). Not all strains are affected in the same way, as can be seen by differences in the amount of inhibition of FAME production when the two different Agr strain pairs are compared (Figs 1 and 2). This strain-dependent difference in Agr control of exoproteins has been described previously [14].

Another difference noted between the two Agr+ strains was the time at which FAME activity was detectable. The ISP479C strain produced detectable FAME activity at 1–2 h, whereas strain RN6390 did not produce detectable levels of FAME until 2–3 h of

Fig. 3. FAME activity of *S. aureus* strains RN6390 (Sar+: ■) and mutant R (Sar−: ●) versus time of incubation and the growth curve of RN6390 (Sar+: □) and RN6911 (Sar−: ○).
growth. This difference does not appear to be due to different growth rates or to differences in optical densities at time zero (Figs 1, 2 and 3).

The Agr" strains began production of FAME early in the stationary phase of growth; however, the rate and amount of FAME produced were much less (Figs 1 and 2). Expression of FAME is not totally eliminated as has been described for class I Agr-regulated extracellular proteins (α-haemolysin, TSST-1, β-haemolysin) [16, 18]. However, the control of FAME production by the agr locus is similar to class II exoproteins (nuclease, β-haemolysin, lipase) in that the protein is still expressed but at much lower levels (50-84% less) [16, 18]. Other Agr class II proteins have shown similar levels of expression in Agr" strains and this is most likely due to the fact that regulation by this locus is leaky [16, 18].

Mutation of the sar locus also affects FAME production. Much less FAME was produced (86% less) in the mutant when compared to the Sar" strain. These results are different from those reported for lipase and protease [14]. Previous studies have shown that sar mutants produce as much or more of these enzymes than the isogenic Sar" strain. This increase in protease may also explain why no late stationary phase increase in FAME activity was noted in the Sar" strain.

Other investigators have provided evidence that the agr locus may be controlled by the sar locus [13]. If a sar mutation occurs then the expression of Agr-regulated proteins will be affected. Much like a toxin, FAME expression is decreased when the sar locus is mutated. However, low levels of FAME would still be expressed due to the fact that Agr does not completely eliminate expression of class II proteins.

Finally, it is hoped to purify FAME, characterise the enzyme and then clone the FAME gene in Escherichia coli; and eventually to produce a strain of S. aureus unable to express FAME and examine the effects of this mutation in a murine abscess model [4].

We thank Drs R. P. Novick, J. J. Iandolo, F. A. Kapral and A. L. Cheung for the use of their bacterial strains. We are grateful to Drs N. Sargentini and G. Tritz for their helpful discussions. We also thank Kathy Krog for excellent computer and typing assistance. This study was partially supported by a Warner fund grant from KCOM and Public Health Service grant R15AI-3119 from the National Institute of Allergy and Infectious Diseases to N.R.C.

References