Protein Antigens Increase the Protective Efficacy of a Capsule-Based Vaccine against *Staphylococcus aureus* in a rat model of osteomyelitis

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Running title: *Staphylococcus aureus* conjugate vaccine

Key words: *Staphylococcus aureus* / vaccine / capsular polysaccharide / osteomyelitis / rat

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SUMMARY

*Staphylococcus aureus* is an invasive bacterial pathogen, and antibiotic resistance has impeded adequate control of infections caused by this microbe. Moreover, efforts to prevent human infections with single component *S. aureus* vaccines have failed. In this study, we evaluated the protective efficacy in rats of vaccines containing both *S. aureus* capsular polysaccharides (CPs) and proteins. The serotypes 5 CP (CP5) and 8 CP (CP8) were conjugated to tetanus toxoid and administered to rats alone or together with domain A of clumping factor A (ClfA) or genetically detoxified alpha toxin (dHla). The vaccines were delivered according to a preventive or a therapeutic regimen, and their protective efficacy was evaluated in a rat model of osteomyelitis. Addition of dHla (but not ClfA) to the CP5 or CP8 vaccine induced reductions in bacterial load and bone morphologic changes compared with immunization with either conjugate vaccine alone. Both the prophylactic and therapeutic regimens were protective. Immunization with dHla together with a pneumococcal conjugate vaccine used as control did not reduce staphylococcal osteomyelitis. The emergence of unencapsulated or small colony variants during infection was negligible and similar for all of the vaccine groups. In conclusion, addition of dHla to a CP5 or CP8 conjugate vaccine enhanced its efficacy against *S. aureus* osteomyelitis, indicating that the inclusion of multiple antigens will likely enhance the efficacy of vaccines against both chronic and acute staphylococcal disease.
INTRODUCTION

Staphylococcus aureus is a medically important opportunistic pathogen that affects individuals in the hospital setting as well as in the community. S. aureus can provoke skin and soft tissue infections, and it can also disseminate to cause invasive life-threatening infections, including septic arthritis and osteomyelitis (1). Osteomyelitis is a progressive infection of the bone marrow and cortex and is frequently caused by S. aureus (2). It is usually preceded by trauma, other nosocomial infections, orthopedic (3) or maxillofacial surgery (4). The control of S. aureus infections in patients with either nosocomial or community-acquired infections has been hampered by the emergence of methicillin-resistant S. aureus (5-7). The high worldwide prevalence of nosocomial MRSA infections was responsible for the intensive use of glycopeptide therapy (8). Although glycopeptides have long been used to treat severe MRSA infections, the increasing prevalence of clinically relevant isolates with reduced susceptibility to vancomycin (9) and the appearance of MRSA resistant to vancomycin (10) have prompted a search for a suitable immunoprophylactic approach to prevent S. aureus infections.

Due to its vast array of virulence factors and the myriad of infection types that it causes, S. aureus presents a unique challenge for vaccine development. A number of S. aureus antigens have been explored as potential vaccine components. Among these, S. aureus capsular polysaccharides (CP), which are anti-phagocytic and key for immune evasion, have been utilized (11). In a phase III clinical trial a conjugate vaccine including CP serotypes 5 (CP5) and 8 (CP8) significantly \( P = 0.02 \) reduced the incidence of S. aureus bacteremia in patients receiving hemodialysis between week 3 and 40 after immunization (12). However, at the study endpoint (week 54) the vaccine efficacy was only 26%, which was not statistically significant. A confirmatory phase III clinical trial failed to reduce bacteremia in hemodialysis patients.
Because of the complexity of *S. aureus* and its myriad of virulence factors, the inclusion of multiple staphylococcal antigens would likely result in a more effective vaccine. Numerous studies suggest that cell wall-linked surface protein clumping factor A (ClfA) is a promising antigen for inclusion in a *S. aureus* multicomponent vaccine. Preclinically, ClfA was shown to be protective in rodent models of arthritis, sepsis and endocarditis (13-15). Alpha toxin (Hla) is a pore-forming exotoxin expressed by *S. aureus* that is cytolytic for a variety of cell types, including platelets, endothelial cells, and monocytes (16). Detoxified Hla induces protection in murine models of lethal pneumonia, subcutaneous abscess formation, and peritonitis (17-19). The next generation *S. aureus* vaccine may benefit from the inclusion of both CPs and protein antigens. The selection of surface antigens for inclusion in an experimental vaccine is difficult because *S. aureus* produces a wide array of surface proteins that promote its virulence but are often redundant in function (20). Moreover, the immune correlates of protection against *S. aureus* infection have not yet been elucidated.

Efforts to prevent staphylococcal osteomyelitis by immunization are few (21). In this study, we evaluated the ability of active immunization to reduce the severity of experimental staphylococcal osteomyelitis, and we compared vaccines that were delivered in a preventive or a therapeutic fashion. CP conjugate vaccines were evaluated alone and in combination with *S. aureus* ClfA or detoxified Hla for their ability to reduce the bacterial burden associated with the disease, as well as to reduce the gross morphologic changes that occur in the bone during chronic staphylococcal infection.
MATERIALS AND METHODS

Bacterial strains. S. aureus clinical strains HU-1 and HU-92a were obtained in 2007 from patients with chronic osteomyelitis at the Hospital de Clínicas José de San Martín, Universidad de Buenos Aires. HU-1 is a CC97 strain that produces CP5, and HU-92a is a CC45 strain that produces CP8. Strains were kept frozen in TSB at -20°C, and S. aureus was routinely cultured at 37°C for 24 h on Columbia agar supplemented with 2% NaCl to enhance CP production. Bacterial cells were harvested and suspended to the appropriate density in trypticase soy broth. S. aureus HU-1 was used as the challenge strain for the CP5 experiments, and S. aureus HU-92a was used for the CP8 experiments.

The rat osteomyelitis model. Outbred Wistar adult rats weighing 280-320 g were purchased from the vivarium of: a) Facultad de Odontología, Universidad de Buenos Aires; b) Facultad de Farmacia y Bioquímica, Universidad de Buenos Aires, or c) Comisión Nacional de Energía Atómica, Ezeiza, Buenos Aires. The rats were housed at the vivarium of the Instituto de Microbiología y Parasitología Médica (IMPaM), Universidad de Buenos Aires and CONICET. Animal care was in accordance with the guidelines set forth by the National Institutes of Health (22). The use of the rat osteomyelitis model was evaluated by the “Comisión Institucional para el Cuidado y Uso de Animales de Laboratorio” (CICUAL), Facultad de Medicina, Universidad de Buenos Aires, and approved through resolution CD 1938/07.

The rats were anesthetized with ketamine/xylazine, the left tibia was exposed, and a hole in the bone was made with a high-speed drill using a 0.4 mm diameter bit. Each tibia was injected with a 5 μl suspension containing ~1x10^6 CFU S. aureus suspended in fibrin glue (Tissucol kit 1 ml, Baxter Argentina-AG Vienna, Austria) (23). Fourteen weeks after bacterial
challenge, the rats were euthanized by CO₂ overdose. Left and right tibias were excised, and adjacent tissue, especially at the distal epiphysis, was removed. The following measurements were made using calipers: a) the distance between the inoculation point and the distal end of the left tibia (DT); b) the left tibia section diameter at the inoculation site (Di) and the perpendicular diameter at the same site (Dp); c) Di and Dp were also measured in the uninfected right tibia of the same rat at the DT determined in the diseased left tibia (control). The osteomyelitic index (OI) was determined as follows: OI=[Dp+Di]<sub>infected</sub> - [Dp+Di]<sub>control</sub>. In pilot experiments we determined that the OI at 7 or 14 wks after injection of sterile Tissucol did not differ significantly from uninjected tibias. Therefore, in order to avoid unnecessary discomfort to rats, only the left tibia was injected, and the OI was determined by subtraction of values from the healthy right tibia (control). One cm segments involving the infected bone were crushed and homogenized in sterile mortars. Homogenates were quantitatively cultured overnight on trypticase soy agar, and the number of CFU was determined. Plates were incubated for 48 h to evaluate the CFU number of S. aureus small-colony variants (SCVs), which were characterized by catalase and coagulase tests and confirmed by a species-specific PCR (24). The OI, the wild-type S. aureus CFU counts and the S. aureus SCV CFU counts from each experimental group (infected and control) were compared. CP5 and CP8 production by S. aureus recovered from the rat tibias was assessed by colony immunoblot as described (25).

**Vaccination schemes.** Rats were vaccinated according to two different schemes. In the preventive scheme, rats were immunized on days 0 and 7. The rats were challenged by the intratibial route with S. aureus on day 28. In the therapeutic scheme, rats were vaccinated 21 d after bacterial challenge and boosted 7 days later (day 28). For both regimens, the rats were euthanized by CO₂ exposure 14 wks (ca. 98 days) after challenge with S. aureus.
Immunogens and controls. Rats were immunized with: Group 1) 10 µg of CP5 or CP8 conjugated to tetanus toxoid [CP5(8)-TT]. Group 2) 10 µg of CP5(8)-TT plus 20 µg of recombinant ClfA region A, domain N123 (ClfA). Group 3) 10 µg of CP5(8)-TT plus 20 µg of purified, genetically detoxified S. aureus α-toxin (dHla). Group 4) 10 µg of Streptococcus pneumoniae CP18C conjugated to TT [Spn-TT]. The 18C-TT conjugate is a pre-GMP lot of one of the conjugates included in a commercial vaccine, and the 18C polysaccharide was fully characterized and devoid of cell wall polysaccharide contamination. Group 5) 10 µg of Spn-TT plus 20 µg ClfA. Group 6) 10 µg of Spn-TT plus 20 µg dHla. Group 7) Phosphate buffered saline (PBS) (control group). The CP/TT ratios were 1/2.6 for the CP5-TT conjugate and 1/2.2 for CP8-TT conjugate. The vaccine antigens (except PBS) were adsorbed on 200 µg Aluminium [(AlPO₄ or Al(OH)₃] and delivered subcutaneously to the rats in a 0.1 ml volume. CP5-TT, CP8-TT, dHla and Spn TT (CP18C) were adsorbed on AlPO₄, whereas ClfA was adsorbed on Al(OH)₃. The selection of AlPO₄ or Al(OH)₃ was based upon intrinsic characteristic of the antigen (such as the isoelectric point) and the pH of the antigen buffer. Preliminary experiments in other settings have shown that the use of AlPO₄ or Al(OH)₃ did not change significantly the adjuvant effect or impact on protection (unpublished data).

Antibody determinations. Blood was collected from all rats by cardiac puncture under anaesthesia prior to euthanasia, and the serum was stored at -20°C. Preliminary experiments showed that the titer of antibodies to CP increased after vaccination, whereas the titer of antibodies in non-vaccinated rats was undetectable and not modified after 14 weeks. Therefore, rats were bled only once at the end of the experiment, and the serum from non-immunized rats served as the control. A similar criterion was adopted for other antibody determinations in order to reduce discomfort and further handling of the animals, as recommended by the CICUAL.
Serum antibody titers to CP5, CP8, Spn CP18C, ClfA and dHla were determined by ELISA using Nunc Maxisorp microtiter plates. Optimal coating conditions were defined for each antigen to insure optimal adsorption on the plates and to allow reproducible detection of specific rat serum antibodies. Plates were coated overnight at 4°C with solutions of purified CP5 (1 µg/ml), CP8 (30 µg/ml), Spn CP18C (40 µg/ml), ClfA (1 µg/ml) or dHla (20 µg/ml) in phosphate buffered saline (PBS). The plates were blocked with PBS + 1% bovine serum albumin (BSA) for 30 min at room temperature with agitation. Two-fold dilutions of rat sera (initial dilution 1:10) were added to the plate and incubated with agitation at room temperature for 30 min. After washing, peroxidase-AffiniPure goat anti-rat IgG, Fcγ fragment specific (Jackson ImmunoResearch Laboratories Inc.; product 112-035-008), diluted 1:5000 in PBS-BSA + 0.2% -Tween 0.05% was added, and the plate was incubated with agitation for 30 min at room temperature. The color was developed using 4 mg o-phenylenediamine + 0.05% H₂O₂ in 0.1 M citrate buffer, pH 4.5, for 15 min in the dark at room temperature. The reaction was stopped with 50 µl 3M HCl and the optical density was recorded at 490 nm relative to 620 nm. The level of antibodies present in the sera was expressed in mid-point titers. Midpoint ELISA titers were calculated by 4-parameter regression analysis and were defined as the reciprocal of the serum dilution that produced an absorbance (at 490 nm) equal to 50% of the maximum value.

**Statistical analysis.** Bone bacterial burdens and OI values were compared by one-way ANOVA, and multiple comparisons were performed with the Bonferroni post-test. Prism 5.0 software (GraphPad) was used for all calculations, and P values <0.05 were considered significant.
RESULTS

CP5 vaccine experiments - preventive scheme. Two experiments of identical design using rats from the same source and vaccinated with preparation from the same lot were performed. Since the data from the individual experiments were similar, the results from identical groups were pooled and presented in Fig. 1. The validity of the experiment was confirmed by the significant correlation of the bacterial load and the OI (r = 0.92; p<0.0007) (Fig. 1C).

The addition of ClfA to CP5-TT slightly reduced the bone bacterial load and the OI compared to that of rats given CP5-TT alone. The differences, however, did not reach were not significant (Figs. 1A and 1B). The addition of ClfA to the Spn-TT conjugate did not improve the protective efficacy of the CP conjugate (Figs 1A and 1B), indicating the limitations of ClfA as a protective immunogen in the osteomyelitis infection model. Nonetheless, the rats given CP5-TT + ClfA show a reduced bacterial burden compared to those animals give Spn-TT + ClfA, suggesting a role for capsular antibodies in reducing infection.

Of note, the addition of dHla to the CP5-TT preparation produced a significant reduction in bacterial load compared to that of CP5-TT alone (Fig. 1A, CP5 vs. CP5+dHla, p<0.001). The specificity of this synergistic response is demonstrated by the fact that dHla did not improve the protective efficacy of Spn-TT (Fig. 1A, Spn vs. Spn+dHla). Rats given the CP5-TT + dHla vaccine showed significantly fewer CFU/bone than the rats given Spn-TT + dHla (p<0.001). The relevance of dHla as vaccine antigen was underscored by the fact that rats given the CP5-TT + dHla showed significantly fewer CFU/bone than the rats given CP5-TT + ClfA (p <0.001).
The greatest reduction in OI occurred in the rats given the CP5-TT + dHla vaccine (Fig. 1B), but the reduction in OI did not reach significance.

**CP5 vaccine experiments - therapeutic scheme.** Two experiments of identical design were performed, and the results were pooled as described above. The overall results of the therapeutic vaccine approach correlated with those obtained with the preventive scheme. The CP-TT conjugates alone showed little effect therapeutically (Fig. 2A). In contrast to the results obtained in the prophylactic regimen, however, the addition of ClfA to the CP5-TT preparation resulted in a modest but significant reduction in the bone bacterial burden compared to that of CP5-TT alone (p<0.05) (Fig. 2A). Similar to the results of the preventive vaccine approach, the therapeutic addition of dHla to the CP5-TT vaccine provided the greatest and most consistent protection against infection compared to CP5-TT alone or Spn-TT + dHla (p<0.001). The greatest reductions in OI (Fig. 2B) occurred in the rats immunized with CP5-TT + dHla, but the differences did not achieve statistical significance. The validity of the experiment was confirmed by the significant correlation of the bacterial load and the OI (r = 0.95; p<0.0009) (Fig. 2C).

**Vaccine antibody levels - CP5 studies.** Serum was obtained from experimental and control groups on the day the rats were euthanized. Serum antibody titers in the rats immunized with the antigens used for the CP5 experiments are shown in Fig 3. Overall, the antibody responses of the immunized rats were specific for the individual immunogens. The CP5-TT antibody response was generally higher than that elicited by the Spn-TT vaccine (Figs. 3A, 3B, 3C, and 3D). Antibody titers resulting from vaccination with either CP conjugate tended to be higher in the rats immunized therapeutically (Fig. 3B and 3D). The overall antibody response to
dHla was greater than that of the other antigens (Fig. 3G and 3H). However, dHla antibody titers in rats immunized with antigens other than dHla were also somewhat higher than those of the heterologous antigens, particularly in the rats immunized therapeutically (Fig. 3H). These data suggest that dHla is an immunodominant protein elicited during staphylococcal infection. Whether protection against infection mediated by the CP5-TT vaccine administered with dHla is mediated by antibodies or by immune T cells remains to be determined.

**CP8 vaccine experiments.** Two experiments each were performed using the preventive and therapeutic vaccination schemes, and the results were pooled as described above. The validity of the CP8 experiments was confirmed by the significant correlation of the bacterial load and the OI (preventive scheme: \( r = 0.97; \ p<0.0001; \) Fig. 4C) (therapeutic scheme: \( r = 0.95; \ p<0.0001 \) ) (Fig. 5C).

The results obtained with the CP8 vaccine groups were similar to those obtained with the CP5 vaccines. In the preventive scheme, vaccination with CP8-TT + dHla significantly reduced the bacterial load \( (p<0.001) \) and the OI \( (p<0.05) \) compared to vaccination with CP8-TT alone (Figs 4A and 4B). Similar findings can be seen for rats immunized therapeutically with either CP8-TT + dHla or CP8-TT alone (Figs. 5A and 5B). In contrast, the addition of ClfA to the CP8-TT vaccine did not improve CP8-TT vaccine efficacy. Of note are the results showing that when dHla was combined with the Spn-TT, no protection against osteomyelitis was observed. Thus, the combination of *S. aureus* CP antigens and dHla provided the best protection against experimental infection with encapsulated staphylococci.
**Vaccine antibody levels – CP8 studies.** Specific serum antibody responses to the various vaccine antigens used for the CP8 experiments are shown in Figure 6. Overall, the antibody responses of the immunized rats were specific for the individual immunogens (Figs. 6A, 6B, 6C, and 6D). Notable differences in antibody titers were not observed in rats immunized with CP-TT conjugates before or after bacterial challenge (Figs. 6A, 6B, 6C and 6D). The antibody responses to dHla in the rats immunized prophylactically (Fig. 6G) were not as striking as those observed in the CP5 experiments (Fig. 3G). Within the CP8 experiment, the most consistent antibody response to Hla was seen in rats immunized therapeutically with dHla given with CP8-TT or Spn-TT (Fig. 6H). Whether protection against infection mediated by the CP8-TT vaccine administered with dHla is mediated by antibodies or by immune T cells is not yet known.

**Emergence of SCVs.** The percent of SCV colonies recovered from the bone cultures of each rat were quantified. Despite the chronic nature of the experimental infection (14 wks), the emergence of SCVs was <1.6% for all of the experimental groups, with a mean recovery of only 0.86%. Statistical analysis by ANOVA revealed no significant differences among any of the rat groups within each of the 4 experiments (CP5-preventive, CP5-therapeutic, CP8-preventive or CP8-therapeutic) (data not shown). Similarly, cultures from only three rats yielded colonies of *S. aureus* that no longer produced CP5 or CP8 (measured by colony immunoblot), and there was no obvious association between experimental group and loss of CP production.
DISCUSSION

A number of *S. aureus* target antigens have been evaluated as vaccine components in preclinical active or passive immunization studies (20, 26-30). Because capsular antigens comprise effective vaccines against other encapsulated bacterial pathogens, early efforts in *S. aureus* vaccine development focused on these components. Covalent coupling of CP5 and CP8 to protein carrier molecules significantly increases their immunogenicity and T-cell dependent properties (31-33). CP5/CP8-based conjugate vaccines alone failed in clinical trials aimed at reducing bacteremia in hemodialysis patients (34, 35). Similarly, a vaccine targeting a single protein (IsdB) failed to reduce surgical wound infections in patients undergoing a median sternotomy (26, 28, 36). Because the nature of *S. aureus* pathogenicity is multifactorial, it is likely that the protective responses required to prevent infection should necessarily target a number of virulence factors (37, 38). The fact that CP-based vaccines induce functional antibodies that mediate opsonophagocytic killing of encapsulated staphylococci supports the inclusion of CP5/CP8 in a multicomponent vaccine. Indeed, CP5 and CP8 conjugates have been included in a recently developed tetravalent vaccine that has entered phase I/II clinical trials (36). In the present study, however, vaccination of rats with the CP5 or CP8 conjugates alone did not significantly diminish the bone bacterial load or the severity of bone infection. The addition of staphylococcal proteins (dHla and to a lesser extent ClfA) significantly reduced the bacterial burden in rats with osteomyelitis.

ClfA is a cell wall-anchored adhesin that mediates *S. aureus* binding to fibrinogen and promotes the attachment of *S. aureus* to biomaterial surfaces, fibrin clots, platelets and damaged endothelial surfaces (39). This surface protein is conserved among *S. aureus* isolates and has shown protective efficacy in diverse animal models (13-15, 40, 41). A pooled human
immunoglobulin G preparation (INH-A21, Veronate®) from healthy donors selected for high antibody titers to *S. aureus* ClfA and *S. epidermidis* SdrG failed to protect neonates from staphylococcal sepsis in a phase III clinical trial (42). Despite the failure of this passive immunotherapy, there is sustained interest in ClfA as a vaccine component for active immunization against *S. aureus* (43). We demonstrated the benefit of targeting both CP5 and ClfA antigens in a mouse mastitis model, in which passive immunization with antibodies to the two antigens had an additive effect on reducing bacterial burden (44). In the present study, the addition of ClfA to the CP5-TT vaccine only reduced the bacterial load in the CP5 therapeutic regimen.

Alpha toxoid was first suggested as a vaccine component in 1977 when it was shown to prevent the lethal gangrenous form of *S. aureus* mastitis in rabbits (45). Subsequently, a number of *S. aureus* vaccines composed of inactivated toxins or their subunits have been evaluated preclinically, and the results of these studies were reviewed recently (20, 26). In addition, *S. aureus* Hla activates the immune system via a TLR2-independent mechanism whereby NOD2 signaling results in protection against a murine staphylococcal infection (46). The fact that a Th17 response is relevant to host defense against *S. aureus* (47, 48) and that Hla is able to induce IL-17A in blood (49) has renewed the interest in Hla as vaccine component. Our results in the osteomyelitis model support the use of dHla in a multicomponent *S. aureus* vaccine. The addition of dHla to either the CP5-TT or CP8-TT conjugate vaccine provided the most striking and consistent protection against the severity of experimental osteomyelitis, as well as the bacterial burden in the infected bone. Whereas vaccination with dHla plus an irrelevant polysaccharide vaccine was not protective, the addition of dHla to CP5(8)-TT induced a significant reduction of the *S. aureus* bone load and the severity of experimental osteomyelitis,
supporting a synergistic effect of the two antigens resulting from immunization. However, only 37% of 76 highly cited animal research studies were replicated in human randomized trials (50). Similarly, a recent study aimed at evaluating how well research from murine clinical models mimics human disease indicates that the transcriptional response to inflammatory stresses in mice is a poor reflection of the human response to similar stresses (51). Thus, data from animal models, although informative, are not necessarily predictive of efficacy in humans (47, 48). Further clinical trials of vaccines that include CP5/8, ClfA and/or dHla are required to ascertain whether the findings of the present investigation translate into success in humans.

Immunization with CP5(8) may result in production of opsonic antibodies that enhance phagocytic clearance of *S. aureus*, whereas immunization with dHla may induce neutralizing antibodies that reduce the tissue damage associated with infection. *S. aureus* is a ubiquitous microbe, and thus it is feasible that the rats were exposed to *S. aureus* prior to experimental infection. This fact may explain the observed elevated baseline antibody levels to Hla in animals not administered dHla. Because all of the animals were subjected to experimental infection, the measurable titers of antibodies to Hla might have resulted from an enhanced response to this antigen by the rats, compared with the other presumably weaker antigens. The immunization schedule utilized was based upon preliminary experiments. An extended immunization schedule might have resulted in a better immune response in the immunized rats. Even though the vaccination schedule might be considered suboptimal, protective efficacy was observed in our studies, supporting the use of multiple antigens in a *S. aureus* vaccine.

SCVs, which have been isolated from patients with chronic, recurrent, or antibiotic-resistant diseases (52), are phenotypes that hold low virulence and that are particularly adapted to the intracellular environment for long-term persistence (53). Once SCVs have emerged, they are
very difficult to eradicate (54). When *S. aureus* was passaged in vivo in the mouse model of mastitis under the selective pressure of antibodies to CP5 or CP8, SCVs were recovered in high numbers from the infected glands. However, SCVs did not emerge when the mice were treated with both CP antibodies and monoclonal antibodies to ClfA (Aurexis ®). In the present study, only a small percentage of SCVs were recovered from the rats with osteomyelitis, irrespective of the vaccine antigen(s) administered. It is likely that the levels of serum antibodies induced by active immunization with the CP5(8) conjugates were insufficient to generate SCVs.

In conclusion, the addition of *S. aureus* protein antigens to a capsular conjugate vaccine enhanced its efficacy in the prevention and therapeutic treatment of experimental *S. aureus* osteomyelitis in rats. ClfA induced only a modest improvement to the efficacy of CP5(8) conjugate, but dHla induced a significant enhancement of conjugate vaccine efficacy. The inclusion of vaccine components designed to neutralize staphylococcal toxins and block adhesins into a multicomponent vaccine preparation may be an effective strategy for the prevention of *S. aureus* infections.

**ACKNOWLEDGEMENTS**

This study was partially supported by GlaxoSmithKline Biologicals S.A. (Rixensart, Belgium), by the Agencia Nacional de Promoción de la Ciencia y la Tecnología, Argentina (ANPCyT PICT 2010-01039) and by the Secretaría de Ciencia y Técnica, Universidad de Buenos Aires, Argentina (UBACyT 200 201 001 003 47). FRB is supported by ANPCyT (PICT 2010-0733). The authors thank Lorena Medina for her dedicated technical assistance.


FIGURE LEGENDS

Figure 1. Protective efficacy *S. aureus* CP5-TT conjugate vaccines administered alone or in combination with ClfA or dHla. Groups of 6-8 rats were immunized according to a preventive scheme (vaccination before bacterial challenge) and challenged by the intratibial route with *S. aureus* HU-1. The effect of immunization on bacterial load (CFU/bone) and OI (mm) are depicted in panels A and B, respectively. Each dot represents determinations made on an individual rat, and the bars represent the geometric means (bacterial load) or the arithmetic means (OI) of each group. Comparisons were performed by one-way ANOVA followed by the Bonferroni post-test, and the *P* values for relevant comparisons are shown. Panel C depicts the correlation of the bacterial load (CFU/bone) and the OI (mm) (Spearman correlation test).

Figure 2. Therapeutic efficacy *S. aureus* CP5-TT conjugate vaccines administered alone or in combination with ClfA or dHla. Groups of 7-9 rats were immunized according to a therapeutic scheme (vaccination after bacterial challenge) and challenged by the intratibial route with *S. aureus* HU-1. The effect of immunization on bacterial load (CFU/bone) and OI (mm) are depicted in panels A and B, respectively. Each dot represents determinations made on an individual rat, and the bars represent the geometric means (bacterial load) or the arithmetic means (OI) of each group. Comparisons were performed by one-way ANOVA followed by the Bonferroni post-test, and *P* values for relevant comparisons are shown. Panel C depicts the correlation of the bacterial load (CFU/bone) and the OI (mm) (Spearman correlation test).

Figure 3. Serum antibody levels to CP5, Spn, ClfA and dHla in groups of immunized rats and controls. Animals were immunized according to a preventive (A, C, E and G) or a therapeutic
(B, D, F, and H) scheme. Blood was collected 14 weeks after bacterial challenge. Each bar
represents the arithmetic mean ± SEM. \( P \) values for selected comparisons were determined by
ANOVA for nonparametric data (Kruskal-Wallis test) followed by the Dunn’s multiple
comparison post-test.

**Figure 4.** Protective efficacy *S. aureus* CP8-TT conjugate vaccines administered alone or in
combination with ClfA or dHla. Groups of 8-9 rats were immunized according to a preventive
scheme (vaccination before bacterial challenge) and challenged by the intratibial route with *S.
aureus* HU-92a. The effect of immunization on bacterial load (CFU/bone) and OI (mm) are
depicted in panels A and B, respectively. Each dot represents determinations made on an
individual rat, and the bars represent the geometric means (bacterial load) or the arithmetic
means (OI) of each group. Comparisons were performed by one-way ANOVA followed by the
Bonferroni post-test, and \( P \) values for relevant comparisons are shown. Panel C depicts the
correlation of the bacterial load (CFU/bone) and the OI (mm) (Spearman correlation test).

**Figure 5.** Therapeutic efficacy *S. aureus* CP8-TT conjugate vaccines administered alone or in
combination with ClfA or dHla. Groups of 8-9 rats were immunized according to a therapeutic
scheme (vaccination after bacterial challenge) and challenged by the intratibial route with *S.
aureus* HU-92a. The effect of immunization on bacterial load (CFU/bone) and OI (mm) are
depicted in panels A and B, respectively. Each dot represents determinations made on individual
rats, and the bars represent the geometric means (bacterial load) or the arithmetic means (OI) of
each group. Comparisons were performed by one-way ANOVA followed by the Bonferroni
post-test, and the $P$ values for relevant comparisons are shown. Panel C depicts the correlation of the bacterial load (CFU/bone) and the OI (mm) (Spearman correlation test).

**Figure 6.** Serum antibody levels to CP8, Spn, CIfA and dHla in groups of immunized rats and controls. Animals were immunized according to a preventive (A, C, E and G) or a therapeutic (B, D, F and H) scheme. Blood was collected 14 weeks after bacterial challenge. Each bar represents the arithmetic mean ± SEM. $P$ values for selected comparisons were determined by ANOVA for nonparametric data (Kruskal-Wallis test) followed by the Dunn’s multiple comparison post-test.
A) Bacterial load (log CFU/tibia)

- PBS
- Spn
- CP5
- CP5+ CifA
- CP5+ dHla
- CP5+ CifA+dHla
- Spn+
- Spn++

Significance:
- p<0.001
- p<0.05
- p<0.001

B) Osteomyelitic index (mm)

- PBS
- Spn
- CP5
- CP5+ CifA
- CP5+ dHla
- CP5+ CifA+dHla
- Spn+
- Spn++

C) Osteomyelitic Index vs Bacterial load (CFU/bone)

- CP5-TT
- Stn-TT
- CP5-TT+CifA
- CP5-TT+dHla
- Stn-TT+CifA
- Stn-TT+dHla
- PBS

- r = 0.95
- p = 0.0009
Antibodies to CP5

CP5 - Therapeutic scheme

**A**
Antibodies to CP5

- **B**: Antibodies to CP5
- **C**: Antibodies to Spn
- **D**: Antibodies to Spn
- **E**: Antibodies to ClfA
- **F**: Antibodies to ClfA
- **G**: Antibodies to dHla
- **H**: Antibodies to dHla

**Summary**

- CP5: **p<0.001**
- ClfA: **p<0.001**
- dHla: **p<0.001**

**Preventive scheme**

- CP5: **p<0.01**, N.S.
- Spn: **p<0.05**
- ClfA: **p<0.01**
- dHla: **p<0.05**

**Therapeutic scheme**

- CP5: **p<0.05**, N.S.
- Spn: **p<0.05**
- ClfA: **p<0.01**, N.S.
- dHla: N.S.
CP8 - Preventive scheme

A) Antibodies to CP8

B) Antibodies to CP8

CP8 - Therapeutic scheme

C) Antibodies to Spn

D) Antibodies to Spn

E) Antibodies to ClfA

F) Antibodies to ClfA

G) Antibodies to dHla

H) Antibodies to dHla