REVIEW ARTICLE

Virulence factors involved in the pathogenesis of bovine intramammary infections due to Staphylococcus aureus

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Summary. Staphylococcus aureus is a major causative agent of intramammary infections in dairy cows. In this report, the pathogenesis of these infections is described. The potential role in virulence of S. aureus surface components (adhesins, protein A and capsular polysaccharides), toxins, extracellular enzymes and coagulase, and perspectives for the development of an efficient vaccine are discussed.

Introduction

Staphylococcus aureus is a gram-positive, facultatively anaerobic, non-sporulating, non-motile, catalase-positive and oxidase-negative coccus. This bacterium, which is a natural inhabitant of mammalian skin and mucous epithelia, is the causative agent of various diseases in man and domestic animals.

In man, S. aureus causes two main types of infection: (1) cutaneous or mucosal infections, and (2) septicemic infections which are generally associated with visceral (abscesses, endocarditis, lung infections) or bone (osteomyelitis) infections.

In domestic animals, S. aureus is mainly involved in intramammary infections (IMI) of lactating females. In cows, IMI due to S. aureus, which account for 25–30% of total IMI, are generally subclinical. Nevertheless, IMI cause considerable economic losses, particularly milk losses which range from 10 to 25% of total yield according to the intensity of inflammation and the stage of lactation when the infection occurs. Moreover, the presence of S. aureus in raw milk used by dairy industries is a public health problem.

Current control of intramammary infections

At present, the control of S. aureus IMI is based on two types of measures. Firstly, hygiene at milking that aims to limit the spread of IMI in herds, particularly pre-milking washing and post-milking drying of udders with individual cloths, dipping teats with antiseptic solutions after milking, and cleaning of milking machines. Secondly, systematic antibiotic treatment by the intramammary route at drying-off to cure chronic subclinical IMI established during lactation and to prevent new infections during the dry period.

In practice, these measures, which are often considered constraining or expensive, are not always applied on the farm. The presently available antibiotic preparations used during lactation cure < 50% of S. aureus IMI. This low efficiency seems not to be due to acquired resistance but may result from the fact that some bacteria are located intracellularly, in macrophages or epithelial cells, and inaccessible to antibiotics.

In the context of the high prevalence and economic consequences of S. aureus IMI, and the relative inefficiency of control measures, the development of a vaccine against S. aureus IMI is of great interest. Numerous attempts at vaccination with live or killed bacterial cells, isolated bacterial cell walls, toxoid or killed cell-toxoid preparations, have been made. In several experiments, particularly those with toxoids, a reduction of the frequency and the severity of clinical IMI has been observed. However, whatever the antigenic preparations used, protection against new infections, which is the main objective of vaccination, has never been achieved. Three major reasons can be invoked to explain these results: (1) an incomplete knowledge of the immune defence mechanisms of the mammary gland; (2) the fact that vaccine prepared in vitro did not contain relevant antigens, i.e., those expressed in vivo and involved in virulence; and (3) the lack of appropriate immunisation schedules.

The development of a vaccine against a bacterial infection requires a clear definition of the major
virulence factors involved in the infection. Molecular genetics permits the isolation of specific genes from bacterial pathogens, the elucidation of their structure and function and the modification of their expression. A powerful approach based on Koch's postulates has been proposed as follows.\(^1\) (1) The supposed virulence factor should be associated with pathogenic strains of the bacterial species under investigation. (2) The gene(s) encoding this supposed virulence factor should be isolated and the inactivation of the gene(s) should lead to a significant loss of virulence. (3) The re-introduction of the gene(s) of origin in the modified strain(s) should restore the virulence. During the past decade, this modern molecular approach has been developed to study virulence factors of \(S.\) \(aureus\) isolates from bovine IMI.

The purpose of this article is to review the current knowledge on the virulence factors involved in the pathogenesis of bovine \(S.\) \(aureus\) IMI and examine the prospects for a vaccine against these infections.

Contamination of the teat orifice and entry of \(S.\) \(aureus\) into the mammary gland

Intramammary infections caused by \(S.\) \(aureus\), and also by other bacterial species, are due to the entry of bacteria through the teat canal into the mammary gland. One quarter of a cow's udder is generally infected independently of the other three.

The main reservoirs of \(S.\) \(aureus\) in herds are infected quarters and the skin of the udder and teats.\(^5,11\) The presence of lesions on teat skin, such as chaps and cracks, allows persistent colonisation of the skin by \(S.\) \(aureus\). A study on c. 800 quarters has shown that without hygienic measures during milking, the rate of teat orifice contamination by \(S.\) \(aureus\) can reach 60% of healthy quarters compared to > 85% of quarters with teat skin lesions.\(^12\) To avoid teat trauma or lesions due to adverse milking conditions, the vacuum level and pulsation frequency of milking machines must be checked regularly and adjusted.

As \(S.\) \(aureus\) is mainly located on or in udders, the transmission of bacteria from teat to teat in the same cow or from one cow to another generally occurs during milking. The main vectors of transmission are the hands of milking staff, the cloths used to wash or dry the udder, the milking cups and contaminated milk which can flow back to the teat during milking as a consequence of sudden variations of vacuum in the milking machine (impact phenomenon).\(^5\) The role of milking in the transmission of \(S.\) \(aureus\) is confirmed by the fact that rigorous hygienic measures during milking lead to a drastic decrease in teat orifice contamination and, consequently, in the rate of new \(S.\) \(aureus\) IMI in herds.\(^13\)

Once the teat orifice is contaminated by \(S.\) \(aureus\), bacteria can persist and multiply\(^14\) and then enter the teat canal either by progressive colonisation or by the impact phenomenon, particularly at the end of milking.\(^5\) Colonisation of the teat canal seems to be a necessary preliminary step for the mammary gland to be infected.\(^15\) \(S.\) \(aureus\), together with coagulase-negative staphylococci, \(Corynebacterium\) \(bovis\) and \(Streptococcus\) \(agalactiae\), are the micro-organisms most frequently isolated from the teat canal,\(^16\) where they can adhere to the keratinised epithelial cells.\(^17,18\) It seems that the closer the bacteria colonising the teat canal are to the teat sinus, the higher is the risk of mammary infection.\(^19\) The contact of these bacteria with milk could allow their transfer into the teat sinus when the intramammary pressure increases (before milking or when the cow walks, lies down or stands up) and just after milking when the teat canal is distended.\(^5,20\)

Colonisation of the mammary gland

Several studies have shown that it is possible to induce experimental \(S.\) \(aureus\) IMI by intramammary infusion of a low number of bacteria (< 100 cfu) through the teat canal.\(^21,22\) These observations demonstrate that milk is a good medium for growth of \(S.\) \(aureus\), which can hydrolyse casein and ferment lactose,\(^1\) and that the natural defence mechanisms of the mammary gland against infection are inefficient. Bacterial surface properties play a major role in the host-bacterium relationship.\(^23\) During \(S.\) \(aureus\) IMI, surface components participate in bacterial adhesion to host mammary tissues and to resistance to phagocytosis by milk cells.

Role of adhesins

The first step in the colonisation of the mammary gland by \(S.\) \(aureus\) seems to be adhesion to epithelial cells. The in-vitro adhesion of \(S.\) \(aureus\) to bovine mammary epithelial cells was demonstrated initially by Frost\(^24\) and then confirmed by other authors.\(^25\) This phenomenon varies with the strain and with the origin of the cells. The adhesive ability of a strain varies little when epithelial cells come from the different quarters of the same cow but greatly when cells come from the udders of different cows.\(^26,27\) The attachment of \(S.\) \(aureus\) to epithelial cells of lactiferous ducts has been confirmed \textit{in vivo} by microscopic examination of sections of experimentally infected mammary glands.\(^28\) The mechanisms of \(S.\) \(aureus\) adhesion to mammary epithelial cells are not well known. Two types of interactions could be involved: (1) non-specific physicochemical interactions, and (2) specific interactions between bacterial cell-wall-associated receptors and host components.

Most \(S.\) \(aureus\) isolates from bovine IMI have a high surface hydrophobicity which can vary from strain to strain and with the culture medium.\(^29,30\) This surface characteristic could favour the fixation of bacteria to host cells through hydrophobic interactions with the cell membrane. Moreover, \(S.\) \(aureus\) can bind to host
extracellular proteins such as collagen \(^{31}\) and fibronectin (Fn). \(^{32}\) In strains of human origin, a relationship between invasiveness (i.e., the ability to travel through the blood stream or tissues to cause disease), adhesion in-vitro systems and quantitative expression of receptors to Fn has been reported. \(^{33}\) The Fn-binding receptor of \(S. aureus\) has been identified as a protein of \(c. 200 \text{ kDa}.\) \(^{34}\) In fact, two different closely linked genes encoding two structurally related Fn-binding proteins (Fnbp) have been recently described and amino-acid homologous units (called D1–D4) have been defined as the Fn-binding domain on these proteins. \(^{35, 36}\) Receptors for collagen and Fn have also been demonstrated on strains isolated from bovine IMI. \(^{37, 38}\) However, the role of Fnbp in the virulence of \(S. aureus\) during IMI has not been definitively demonstrated. Molecular biology could provide the tools to clarify this point. As the complete sequences of the genes encoding Fnbp are well established, \(^{39, 40, 41}\) it will be possible to isolate mutants lacking Fn-binding ability by site-specific mutagenesis. These mutants could be used in experiments such as in-vitro adhesion to mammary cells and experimental IMI in cows. It is noteworthy that the surface components of \(S. aureus\) that mediated adhesion either by hydrophobic interactions or by specific recognition of host components are proteins. Adherence of \(S. aureus\) to ductular cells \(\text{in vitro}\) is inhibited when bacteria are first treated with proteolytic enzymes. \(^{39}\) These enzymes also decreased surface hydrophobicity \(^{39}\) and the ability of \(S. aureus\) to bind collagen \(^{31}\) and Fn. \(^{37}\) Recently, Lindahl \(\text{et al.}\) \(^{46}\) purified a cell-wall associated protein of 145 kDa from a strain of \(S. aureus\) isolated from bovine IMI. This component has the ability to bind to membrane proteins of bovine milk fat globules and mammary epithelial cells.

From facts already known, it can be supposed that adhesins such as Fnbp could mediate \(S. aureus\) adhesion to epithelial cells and to micro-lesions of the mammary epithelium where basal lamina and inflammatory conjunctive tissue, which are rich in Fn and collagen, are exposed. \(^{37}\) In the early stages of the infection, adhesion to epithelial cells could prevent \(S. aureus\) from flowing out of the gland during milking. Then, \(S. aureus\) could disseminate in the mammary gland by adhering to rising fat globules \(^{17, 41}\) and then attach to the epithelial cells at different levels of the gland, i.e., teat sinus and lactiferous sinuses and ducts. \(^{26}\)

**Expression of antiphagocytic factors by \(S. aureus\)**

The main defence mechanism of the mammary gland against infections by \(S. aureus\) is phagocytosis by polymorphonuclear neutrophil leucocytes (PMNL). \(^{42}\) Schalm \(\text{et al.}\) \(^{43}\) have demonstrated that the induction of neutropaenia by continuous intravenous injection of equine anti-bovine leucocyte serum led to a conversion of \(S. aureus\) IMI from subclinical to the acute gangrenous form. Moreover, Postle \(\text{et al.}\) \(^{28}\) failed to induce experimental \(S. aureus\) IMI in quarters with milk somatic cell counts (SCC) greater than \(6 \times 10^9\) cells/ml.

The SCC of milk from healthy quarters is generally \(< 10^4\) cells/ml. Infection of the mammary gland entails a flow of leucocytes, mostly PMNL, from blood through the mammary parenchyma into milk. \(^{44}\) After experimental intramammary inoculation with \(S. aureus\), the SCC of milk begins to increase \(1-3\) h after inoculation and can reach several million cells/ml within \(24-48\) h. \(^{28, 45}\)

Phagocytosis of bacteria by PMNL includes recognition, ingestion and intracellular destruction of bacteria. \(^{46}\) Membrane receptors of PMNL can bind non-specifically to bacterial surface components through lectin-carbohydrate-like interactions but recognition of bacteria and subsequent activation of PMNL are much more efficient when they are mediated by opsonins. Opsonisation of \(S. aureus\) involves antibodies, particularly those specific to peptidoglycan epitopes, and the C3b component of complement. \(^{47}\) The C3b component is generated by activation of complement by the classic or alternate pathways, either directly by surface components of \(S. aureus\) such as peptidoglycan or teichoic acid, \(^{48}\) or by immune complexes between peptidoglycan and specific antibodies, particularly of the IgG isotype. \(^{47}\) Fixation of immunoglobulins (associated to C3b or not) by their antibody site on bacteria and by their Fc fragment on PMNL surface receptors activates the ingestion of bacteria by PMNL. In ruminants, most of the membrane Fc receptors on PMNL are specific to IgG2 isotype. \(^{49, 50}\)

The milk of a healthy quarter has a low concentration of opsonins, complement \(^{51}\) and immunoglobulins (0.8–1 mg/ml, i.e., 25–30 times less than in serum). \(^{52}\) The milk content of IgG2, which is the major isotype involved in opsonisation, is particularly low (0.05–0.06 mg/ml). The inflammation of the mammary gland during \(S. aureus\) IMI includes both PMNL recruitment and exudation of plasma through the mammary parenchyma into milk. Milk concentrations of IgG1, IgG2 and complement increase. \(^{53, 54}\) Nevertheless, this inflammation does not result in the total elimination of bacteria but only in a partial control of the infection which evolves to a chronic subclinical form that reflects a balance between multiplication of bacteria and the mammary defences. This balance can be explained on one hand by the low efficiency of phagocytosis due to the ingestion of casein and fat globules by PMNL \(^{55, 56}\) and to the low level in milk of opsonising antibodies specific to \(S. aureus\) and, on the other hand, by the production of antiphagocytic factors such as protein A and a capsule by \(S. aureus\).

**Protein A.** Staphylococcal protein A (SpA) is a protein bound to the surface of the bacterial cell wall that has the ability to bind IgG of numerous mammalian species (man, mouse, rabbit, ruminants) by the Fc fragment \(^{57, 59}\). Using human IgG as a source of opsonins, Dossett \(\text{et al.}\) \(^{60}\) have demonstrated that bacterial bound or free
soluble SpA inhibited phagocytosis of *S. aureus* by human PMNL *in vitro*. Nevertheless, this anti-phagocytic effect seems to depend on the relative availability of both antibodies and complement. In the presence of human IgG-deficient serum, opsonisation of SpA high-producing strains is more efficient than that of low-producing strains whereas the converse has been observed with normal human serum or purified IgG as opsonising.  

SpA, which has been detected on 50–60% of strains isolated from bovine IMI, can react strongly with bovine IgG2, but weakly with IgG1. In milk from a healthy quarter, the concentration of IgG2, the major isotype involved in opsonisation, is very low. Consequently, SpA may play a minor role in *S. aureus* virulence, at least during the early stages of bovine IMI. Postle *et al.* reported a highly significant negative correlation between the amount of SpA on *S. aureus* strains (12 strains studied) and their experimental pathogenicity in cows. This was postulated to be due to complement activation by SpA that released chemotactic (C3a and C5a components) and opsonising (C3b) factors. Conversely, during later stages of IMI when the opsonising IgG2 milk content increases as a consequence of plasma inflammatory exudation, the antiphagocytic effect of SpA could be expressed. Nevertheless, this hypothesis has still to be proved. Finally, it is noteworthy that Jonsson *et al.* observed no difference in virulence for the mouse mammary gland between an *S. aureus* low-producing strain, a high-producing mutant and a SpA-negative mutant carrying a plasmid containing the complete gene for SpA. With the same mouse model, Foster *et al.* observed that SpA-deficient mutants were less virulent than a parental strain that produced a high amount of SpA. The discrepancy between these two studies could be due to the method used to isolate mutants—chemical mutagenesis, that can induce pleiotropy in the former study and site-specific mutagenesis in the latter one.

**Capsule.** The capsule is defined as a polysaccharide layer covering the cell wall. When observed by light or electronmicroscopy in the presence of Indian ink, it appears as a white halo surrounding the bacteria. Capsulate *S. aureus* strains produce diffuse colonies in serum-soft agar (SSA), a medium containing normal rabbit serum 1%, whereas non-capsulate strains produce compact colonies. Two studies (each with about 800 strains) have demonstrated that only 5% of *S. aureus* strains isolated from human infections produce a capsule as defined by the microscopic Indian ink and the SSA technique.

The existence of a capsule on *S. aureus* strains isolated from bovine IMI is a much debated matter. Several studies have shown that most strains (85–95%) produce diffuse colonies when inoculated directly from infected milk into SSA. In these studies, SSA was prepared from a modified staphylococcus medium 110 (mod 110) which is enriched with mannitol, lactose and sodium chloride and is known to enhance the production of extracellular polysaccharides by *S. aureus*. Nevertheless, the reliability of the SSA technique for demonstrating capsulation has been contested. On strains isolated from IMI that produced diffuse colonies in SSA, Anderson and Rather *et al.* failed to observe a capsule by light microscopy. They concluded that strains from bovine IMI were not capsule- or that these strains do not produce capsule but slime, a polysaccharide component loosely bound to the bacterial surface. 

Watson’s team in Australia has observed that *S. aureus* strains isolated from bovine or ovine IMI produce a surface component when cultivated *in vitro* in the peritoneal cavity of sheep but not *in vivo* in classical media, which they called a pseudocapsule. The pseudocapsule is associated with the bacterial surface but is not visible by light microscopy and, therefore, cannot be classified as a classical capsule. Watson and Watson demonstrated by electronmicroscopy that the pseudocapsule is also produced *in vivo* during IMI and *in vitro* when *S. aureus* is cultivated in nutrient broth containing bovine, ovine or caprine milk whey. However, this pseudocapsule has never been purified and its biochemical nature and structure have never been demonstrated. Therefore, it cannot be related to the well defined capsular polysaccharides described by Karakawa and Vann (see below). Strains producing a pseudocapsule are more virulent for mice and have a higher resistance to phagocytosis by PMNL. Vaccination of cows or ewes with a pseudocapsule-producing strain induced serum pseudocapsular antibodies and increased the resistance of vaccinated animals to experimental infections by *S. aureus*, as assessed by the severity of clinical signs and milk production. However, pseudocapsular antigens used to detect antibodies induced by vaccination were prepared by abrasion of the vaccine strain surface with glass beads and removal of SpA by elution through an affinity column of Sepharose 4B to which purified rabbit IgG was coupled. It is likely that the extract is a complex multi-antigenic mixture and, consequently, the partial protection observed cannot be ascribed to any single factor.

It has been demonstrated that *S. aureus* strains isolated from human infections produce capsular polysaccharides (CP) that belong to 11 different serotypes. Several serological surveys with rabbit polyclonal sera or mouse monoclonal antibodies (MABS) have shown that 70–80% of strains from human sources belong to two serotypes, 5 and 8. Types 5 and 8 CP have been purified, visualised on bacterial surfaces by electronmicroscopy with gold-labelled MABS and their chemical structures have been defined. They are heteropolymers of N-acetylfucosamine and N-acetylmannuronic acid with O-acetyl groups, the location of which varies according to serotype. These CP do not constitute classical capsules, i.e., visible by light microscopy, but microcapsules. Strains producing CP are resistant to phago-
cytosis in vitro by human PMNL in the presence of heat-inactivated normal human serum but this resistance is suppressed in the presence of CP-specific rabbit serum or mouse MAb.60 By using anti-type 5 and 8 MAb, we have demonstrated recently that these CP types account for > 50% and for c. 20% of 212 S. aureus strains isolated from bovine IMI, respectively.61 These two CP types are also prevalent amongst S. aureus strains from caprine and ovine IMI.61 Most CP-producing strains exhibited diffuse colonies in mod 110-SSA with a masking of cell-wall-associated components such as SpA and teichoic acid.62 Nevertheless, no capsule was observed by the microscopic Indian ink technique. It seems that most S. aureus strains from IMI express a capsule that is sufficiently thick to mask bacterial surface antigen but not thick enough to be detected by light microscopy.63 After cultivation on bovine milk agar, S. aureus strains isolated from IMI become more resistant to phagocytosis in vitro by bovine PMNL in the presence of normal, i.e., complement-containing, bovine serum than after cultivation on conventional media. This resistance is also associated with cell-surface masking ascribable to CP expression, as demonstrated by agglutination with specific MAb.64 With an enzyme-linked immunosorbent assay (ELISA), we have shown that normal bovine serum contains large amounts of anti-teichoic acid antibodies but very low amounts of anti-CP antibodies.64 Consequently, it can be concluded either that CP could act as a barrier to the binding of serum opsonins to S. aureus surface,65 or, if the capsule is permeable to opsonins, as suggested previously,66 CP could prevent their interaction with PMNL-specific membrane receptors.67 Our results are not consistent with those obtained by Xu et al.68 These authors have shown that type 5 and 8 micro-capsule strains did not resist phagocytosis by human PMNL in the presence of normal human serum. These discrepancies could be explained by the fact that our experiments64 were performed with strains of bovine origin cultivated in the presence of milk that enhanced the expression of CP, as demonstrated by the shielding of bacterial surface, whereas Xu et al.67 used strains from man cultivated in conventional Columbia broth. In the latter case, it can be supposed that the microcapsule was not thick enough to impede the interaction between cell-wall associated C3b and receptors to C3b on PMNL.

Hypothetic kinetics of in-vivo expression of S. aureus surface components

Although the kinetics of expression of surface components have never been investigated, it can be supposed that during the first stages of infection S. aureus produces hydrophobic surface proteins, particularly receptors for host components, that allow it to adhere to mammary epithelium and milk fat globules and thus to disseminate in the gland.69,70 Then, during exposure to milk, S. aureus could express CP that prevents phagocytosis by impairing interactions between bacteria and PMNL mediated by opsonins as described above,67 or because of their hydrophilic nature.68 Expression of capsule by S. aureus during IMI has been demonstrated clearly by detection of CP by ELISA with MAb in milk from infected mammary glands69 and by electron-microscopy studies of strains separated from IMI milk samples by magnetic particles coated with anti-capsule S. aureus antibodies.100 These hypotheses are supported by the following observations. The binding of Fn and collagen has been found to be lower for S. aureus grown in milk whey than for those grown in conventional medium.70 Also, growth of S. aureus on milk agar masked cell wall-associated components (SpA and teichoic acid) and increased expression of CP.71 Few data are available on the regulation of the expression of surface or extracellular components by S. aureus. However, it has been demonstrated that the synthesis of a number of exoproteins including α, β and δ toxins, toxic shock syndrome toxin 1 (TSST-1), staphylokinase and SpA was controlled by a locus on the chromosome that was named agr (for accessory gene regulator) and that acts on the transcription of mRNA.101 The existence of a regulatory mechanism of the expression of CP and surface receptors such as Fnbp has yet to be demonstrated.

Pathological modifications to the mammary gland

S. aureus IMI often start with an acute phase with general (fever and anorexia) and local (congestion and hardness of the udder, milk clots) clinical signs. Thereafter, the infection generally becomes chronic and subclinical with sporadic clinical events. Acute gangrenous IMI characterised by a rapid and massive multiplication of bacteria and a general necrosis of the infected quarter is uncommon in cows but frequent in goats and ewes.50 The pathological changes of the mammary gland during S. aureus infection have been investigated by microscopic examination of sections of mammary glands either naturally infected102,103 or experimentally inoculated with bacteria through the teat canal104,105 or into the parenchyma through the skin.50 Different histological lesions have been reported. (1) Early ulceration and erosion of lactiferous sinus and ductular epithelia. (2) Infiltration of the parenchyma conjunctive tissue by macrophages and PMNL which cross the mammary epithelium into lactiferous ducts and glandular alveoli. (3) Lesions of the alveolar secretory epithelial cells of variable severity, from a simple decrease of secretory activity to a total cell lysis. (4) Shrinkage of alveoli, proliferation of conjunctive tissue, accumulation of cellular fragments in glandular alveoli as a consequence of the lysis of epithelial cells and PMNL. These phenomena can lead to the oc-
clusion of alveoli and the trapping of bacteria. The resulting focus of infection can then release bacteria that infect other areas of the gland or can evolve to a granuloma. An ultrastructural study of the mammary glands of mice experimentally inoculated with *S. aureus* has shown that bacteria can be located in the cytoplasm of epithelial secretory cells. It would be interesting to know if bovine mammary epithelial cells also possess the ability to phagocytose bacteria, as has been shown for bovine endothelial cells. This could be studied by using cultured mammary cells and if such is the case, the role of adhesins in the initial adherence of bacteria to cell membrane and of toxins in the subsequent cell lysis must be clarified.

Bacteria are not always present on injured epithelium or tissue. It seems that toxins and enzymes produced by *S. aureus* can diffuse in milk and act at a distance from their site of elaboration. Moreover, lysosomal enzymes released by lysed PMNL could contribute to the alteration of mammary glandular epithelium.

Anderson stressed the dynamic nature of staphylococcal IMI. Pathological modifications affect variable areas of the gland with variable degrees of severity. Conversion of the infection from subclinical to clinical form could reflect the spreading of bacteria through the gland, which depends on the efficiency of local defence mechanisms.

**Role of toxins**

*S. aureus* can produce four different haemolytic toxins: α, β, γ and δ toxins. Two, α and β toxins, appear to play a major role in the virulence of *S. aureus*. α Toxin, which is produced by between 20% and c. 50% (Poutrel, unpublished data) of strains from bovine IMI, is a cytolyisin that binds to cell membranes and forms hexameric pores leading to cell death as a consequence of rapid egress of cytoplasmic components. β Toxin, a type C sphingomyelinase, is produced by 75–100% of these strains.

Early studies have shown that systemic vaccination of goats, ewes and cows with a mixture of α and β anatoxins induced serum antibodies specific to α and β toxins. Vaccination with anatoxins does not lead to the elimination of experimental *S. aureus* IMI but to a decrease in clinical severity, particularly in small ruminants. This effect can be ascribed to toxin neutralisation by antitoxin antibodies which flow from blood to milk during mammary inflammation. Le Gall and Plommet have observed that the severity of experimental *S. aureus* IMI in non-vaccinated ewes is inversely proportional to blood antitoxin antibody titre. Similarly, the blood antitoxin titre in cows increases with age, whereas the incidence of clinical *S. aureus* IMI decreases.

Several recent studies with experimental inoculations of rabbits and mice have confirmed the role of α and β toxins in *S. aureus* IMI. Injection of purified α toxin into rabbit mammary gland caused haemorrhagic necrosis of the gland whose extent depended on the dose, whereas injection of purified β toxin induced inflammation of the gland, i.e., oedema and a flow of PMNL into mammary ducts and glandular alveoli. Simultaneous injection of both toxins had the same effect as that of purified α toxin alone. In the rabbit two different forms of *S. aureus* IMI occur: a chronic form with mammary abscesses and a gangrenous form which is often lethal. Adlam et al. infected the mammary glands of rabbits with bovine mastitis organisms and showed that bacteria can be located in the glandular alveoli and that strain CN6708 probably produced a small amount of α toxin. The roles of α and β toxins in virulence have been investigated in a mouse model with *S. aureus* mutants. Several studies have shown that mutant strains altered in α toxin expression by chemical mutagenesis were less virulent for mouse mammary gland than the α toxin-producing parental strain. From strains producing α and β toxins, Bramley et al. isolated strains defective in α toxin (non-α), β toxin (non-β) or in both α and β toxins (non-αβ) strains expression by directed mutagenesis. The parental and mutant strains were injected into mouse mammary glands to study bacterial growth and pathological lesions *in vivo*. It was concluded that necrosis of the mammary gland and the high death rate in mice (60%) following inoculation of the parental strain was due to α toxin. Both α and β toxins promoted bacterial growth in the mammary gland. However, β toxin appear to contribute little to the pathogenesis of acute mastitis in the mouse model. In the case of non-αβ strains, PMNL and cellular architecture of glandular alveoli were not altered, whereas non-β strains caused PMNL lysis, destruction of the secretory epithelium and foci of necrosis. After induction of inflammation by intramammary injection of *Escherichia coli* lipopolysaccharide, bacterial inoculation caused chronic IMI with all strains, irrespective of toxin expression. Lipopolysaccharide caused an early recruitment of PMNL, which reduced bacterial growth and survival and, therefore, toxin production in milk. This situation, already described, could represent the natural chronic *S. aureus* IMI commonly observed in cows.

Most *S. aureus* isolates from bovine milk samples...
produced leucocidin under in-vitro conditions.121 This toxin, which consists of two separate and synergetic components F and S,122 has cytolytic effects on bovine PMNL.123 Moreover, it has been shown that cows with chronic S. aureus IMI have significant anti-leucocidin antibody titres in serum and milk, compared with non-infected animals.124 The recent cloning and sequencing of leucocidin125 should allow site-specific mutagenesis of the corresponding gene to demonstrate the actual role of leucocidin in the virulence of S. aureus during IMI.

A recent study on 262 S. aureus isolates from bovine IMI has shown that ca. 20% of isolates produced TSST-1, generally in association with enterotoxins C and D.126 TSST-1 is known to induce the release of inflammatory factors such as γ-interferon and interleukin-1 by human mononuclear cells.127,128 Although it is obvious that IMI can be caused by strains which do not produce TSST-1, the role of this toxin has yet to be evaluated.

Role of extracellular enzymes and coagulase

S. aureus produces numerous extracellular enzymes129 including hyaluronidase, phosphatase, nuclease, lipase, catalase, staphylokinase and proteases, that have been implicated in the pathogenesis of bovine IMI. The ability of S. aureus to multiply rapidly in milk is probably a major component of its virulence. Enzymes could allow S. aureus to use milk substrates for their metabolism and, consequently, to adapt to milk and grow.130 However, the involvement of staphyloccocal enzymes in the alteration of mammary tissues remains to be demonstrated clearly.

The role of coagulase in the virulence of S. aureus has been mostly studied in the mouse. Injection of purified coagulase into the mouse mammary gland caused a PMNL flow into secretory alveoli and the involvement of staphylococcal enzymes in the demonstrated clearly. Enzymes could allow S. aureus to use milk substrates for their metabolism and, consequently, to adapt to milk and grow.130 However, the involvement of staphyloccocal enzymes in the alteration of mammary tissues remains to be demonstrated clearly.

The multiplication of S. aureus in milk before the onset of inflammation is undoubtedly of great importance for the establishment of the infection. It is crucial to obtain a recruitment of activated PMNL into milk of vaccinated animals as early as possible after the entry of S. aureus into the mammary gland. This recruitment does not need to be massive as it is likely that a milk SCC of ca. 10^6 cells/ml is sufficient to impair staphyloccocal IMI,131 but it must coincide with the presence of opsonising antibodies in milk. The immune-mediated recruitment of PMNL in the bovine mammary gland has been demonstrated clearly by De Cueninck.134 In cows sensitised to ovalbumin by subcutaneous injection of this protein during lactation, intramammary infusion of ovalbumin elicited inflammation, i.e., release of PMNL into milk. Such a

Conclusions: perspectives of vaccine

During the past two decades, the mouse mammary gland model has been used extensively to study the virulence factors involved in the pathogenesis of S. aureus IMI. This model offers the opportunity to use a large number of animals and given the small size of the glands, to study bacterial growth and the histopathological modifications of the gland in vivo. However, it is likely that results obtained with this model are not fully relevant to natural bovine IMI and, thus, they must always be confirmed by experimental infections of cow udders.

The purpose of vaccination is to increase the efficiency of the natural defences of the mammary gland against S. aureus infection. The ideal vaccine must satisfy the following conditions.9 (1) It must induce the appropriate type of immune response. (2) Antigens that lead to undesirable side effects must be excluded from the vaccine, particularly those inducing severe inflammation, i.e., delayed hypersensitivity, which can cause damage to secretory tissues and thus compromise milk production. With vaccines containing toxoids and killed pseudocapsule-producing strains, Watson135 has shown recently that, following intramammary challenge, vaccinated heifers were more resistant to clinical mastitis and had greater milk production than controls. However, we are convinced that control of immune response and safety of the vaccine could be more easily reached by using selected purified and well-characterised bacterial antigens instead of crude whole-cell preparations.

There is evidence that three major factors could be involved in S. aureus virulence: surface protein receptors to host components such as Fnbp, CP and toxins, particularly α and β toxins. Induction in cows of antibodies against these components could lead to several protective effects: inhibition of S. aureus adhesion to mammary epithelial cells, suppression of resistance to phagocytosis by PMNL and decrease of tissue lesions by neutralisation of toxins. However, the use in vaccination of surface components not involved in virulence but which are potential targets for opsonising antibodies must not be ruled out.

The multiplication of S. aureus in milk before the onset of inflammation is undoubtedly of great importance for the establishment of the infection. It is crucial to obtain a recruitment of activated PMNL into milk of vaccinated animals as early as possible after the entry of S. aureus into the mammary gland. This recruitment does not need to be massive as it is likely that a milk SCC of ca. 10^6 cells/ml is sufficient to impair staphyloccocal IMI,131 but it must coincide with the presence of opsonising antibodies in milk. The immune-mediated recruitment of PMNL in the bovine mammary gland has been demonstrated clearly by De Cueninck.134 In cows sensitised to ovalbumin by subcutaneous injection of this protein during lactation, intramammary infusion of ovalbumin elicited inflammation, i.e., release of PMNL into milk. Such a
recruitment was not observed in control non-immunised animals. With a similar experiment in the guinea-pig, De Cueninck has shown that this phenomenon could be transferred by peritoneal cells but not by serum from immunised animals to control animals. Similarly, Rainard et al. have recently immunised cows with purified protein X, a surface antigen of *Str. agalactiae*. The inflammatory response of the mammary gland, i.e., PMNL recruitment and bactericidal activity of milk, to an experimental infection by a protein X-bearing strain was earlier and much higher in immunised cows than in control cows. Taken together, these results suggest that PMNL functions, including the recruitment into the lumen of the infected mammary gland and the ability to kill invading bacteria, could be increased by immunocompetent cells, presumably T lymphocytes, sensitised by vaccination with bacterial protein antigens. Thus, it would be of great interest to determine if immunisation of cows with *S. aureus* antigens such as Fnbp, anatoxins and CP could induce the recruitment of activated PMNL in the mammary gland during infection by this bacterium.

However, *S. aureus* CP, which do not carry T cell epitopes, are not good immunogens in cows. By contrast, conjugation of CP to a protein carrier, *Pseudomonas aeruginosa* exotoxin A or diphtheria toxoid, increased their immunogenicity and T cell dependence in mice. Similar results have been obtained recently in cows with a *S. aureus* type 5 CP-ovalbumin conjugate. Furthermore, immunisation of mice led to antibodies that promoted opsonisation of *S. aureus* by human PMNL.

Consequently, a vaccine containing purified surface proteins involved in *S. aureus* adhesion (e.g., Fnbp), α and β anatoxins and CP belonging to predominant serotypes and conjugated to one or several of these proteins could be used to immunise cows. It is noteworthy that a recent preliminary study has shown that vaccination of cows with a fusion protein containing the Fn-binding domain of Fnbp lead to significant protection against experimental *S. aureus* IMI, both in terms of clinical signs and reduced SCC. Several parameters will need to be investigated to guarantee the efficiency of the proposed conjugate vaccine—the conjugation technique, the route and timing of administration and the adjuvant. Once these parameters of vaccination have been optimised, the efficient protection of cows against *S. aureus* IMI might be feasible.

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