Anti-staphylococcal activities of lysostaphin and LytM catalytic domain

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Abstract

Background: Lysostaphin and the catalytic domain of LytM cleave pentaglycine crossbridges of Staphylococcus aureus peptidoglycan. The bacteriocin lysostaphin is secreted by Staphylococcus simulans biovar staphylolyticus and directed against the cell walls of competing S. aureus. LytM is produced by S. aureus as a latent autolysin and can be activated in vitro by the removal of an N-terminal domain and occluding region.

Results: We compared the efficacies of the lysostaphin and LytM catalytic domains using a newly developed model of chronic S. aureus infected eczema. Lysostaphin was effective, like in other models. In contrast, LytM was not significantly better than control. The different treatment outcomes could be correlated with in vitro properties of the proteins, including proteolytic stability, affinity to cell wall components other than peptidoglycan, and sensitivity to the ionic milieu.

Conclusions: Although lysostaphin and LytM cleave the same peptide bond in the peptidoglycan, the two enzymes have very different environmental requirements what is reflected in their contrasting performance in mouse eczema model.

Background

The problem of growing antibiotic resistance has been solved only in part by the introduction or reintroduction of new antibiotics (such as the quinupristin/dalfopristin Synercid [1] and the oxazolidinones [2]). Peptidoglycan hydrolases represent an alternative to small molecule antibacterials, despite concerns relating to immunogenicity, the release of proinflammatory components during bacteriolysis and the development of resistance [3]. The peptidoglycan endopeptidases lysostaphin and LytM cleave the characteristic pentaglycine crossbridges of S. aureus peptidoglycan [4-6] and are therefore of interest as potential antistaphylococcal agents.

Lysostaphin (Figure 1) is produced by Staphylococcus simulans biovar staphylolyticus. The secreted preproprotein is synthesized with a leader sequence, proregion, catalytic domain, and the cell wall targeting domain (CWT) [7]. The low complexity proregion consists of a variable number of stereotypical repeats (sequence AEVETSKAPVENT)[8]. It can be cleaved off in vivo by extracellular cysteine protease [9] to release the mature form, which is often simply called lysostaphin and is commercially available. Mature lysostaphin consists of the catalytic and CWT domains. The catalytic domain belongs to MEROPS family 23 in clan MO [10] and can be classified with the LAS metallopeptidases [11]. Sequence alignments suggest that the single Zn2+ ion in the active site is coordinated by His279, Asp283 and His362 (numbering according to Swiss-Prot entry P10547) and a water molecule. As the name implies, the CWT domain anchors the protein to cell walls [9] (Figure 1).

The biological role of lysostaphin is well established. The (mature) protein is inactive against the producer organism, but very effective in cleaving S. aureus cell walls [14]. This property has made the enzyme attractive as an antibacterial agent [15-21]. The protein has been applied to disrupt S. aureus and S. epidermidis biofilms on artificial surfaces [22] and has also been tested as a coating for catheters [23]. In a mouse model, lysostaphin has been used to eradicate S. aureus biofilms from a catheterized jugular vein [24] and also for treatment of systemic
infections [25]. In a cotton rat model, a lysostaphin cream has proven effective in eradicating \textit{S. aureus} nasal colonization [26]. In humans, lysostaphin has been used on an experimental basis to treat methicillin-resistant \textit{S. aureus} aortic valve endocarditis [27]. As the elimination of \textit{S. aureus} carriage in hospital staff is demonstrably effective in reducing infection rates in surgical patients and those on hemodialysis [28], a lysostaphin cream to treat infected, but asymptomatic hospital staff, has potential.

\textit{Staphylococcus aureus} LytM (Figure 1) is an autolysin under the control of the two-component system WalKR,
which is thought to play a role in virulence and cell wall metabolism [29]. The protein is synthesized with a signal peptide (LytM1-25), followed by an N-terminal domain that is homologous to the staphylococcal secretory antigen A (SsaA), another WalKR controlled protein, but not to the N-terminal domain of lysostaphin. The C-terminal domain of LytM can be divided into an occluding region and a region of high similarity to the lysostaphin catalytic domain (52% amino acid identity over 106 residues). The lysostaphin active site residues are all conserved, with a central Zn$^{2+}$ ion that is coordinated by His210, Asp214 and His293 of the catalytic domain [12]. Nevertheless, the structure strongly suggests that full length LytM cannot have significant activity, because the active site is occluded. The expected water molecule in the coordination sphere of the Zn$^{2+}$ ion is displaced by an “asparagine switch” residue (Asn117) of the occluding region, which also blocks part of the active site cleft [12]. However, the crystal structure suggested that the catalytic domain alone should be more active than the full length protein. This was confirmed for a tryptic fragment (LytM180-316, previously referred to as in vitro activated LytM) and for the recombinantly overexpressed catalytic domain (LytM185-316, previously referred to as active LytM) [12,30]. In this work, we use the designation “catalytic domain” for the LytM185-316 fragment for consistency with the well-established lysostaphin nomenclature, even though the catalytic domain and occluding loop form the globular unit in the full length protein [12]. LytM lacks a counterpart for the cell wall targeting domain of lysostaphin (Figure 1).

The biological role of LytM is still not clear [31]. The protein was originally described as an autolysin (detected in an otherwise autolysin deficient background) [5] and reported to have glycylglycine endopeptidase activity [32]. Both we and other investigators have since then carried out experiments that cast doubt on the peptidoglycan hydrolyzing activity of full length LytM [12,31]. Nevertheless, the data clearly confirmed such activity of the catalytic fragment [12,30]. It remains to be determined whether the LytM catalytic domain can be released under physiological circumstances. A proteomic study of the S. aureus cell wall envelope fraction has identified only full length LytM (with a molecular mass of approximately 40 kDa and a pI around 6), but not in the predicted active form [33]. Although the physiological role of LytM and its catalytic domain remains uncertain, the catalytic domain has properties that could make it attractive as a potential antistaphylococcal agent. First, the protein can be easily overexpressed in Escherichia coli with very high yields and is easy to purify [30]. Moreover, preliminary in vitro experiments indicated that in certain conditions LytM185-316 was similarly effective as lysostaphin in clearing turbid cell wall suspensions.

Therefore, we proceeded to compare lysostaphin and LytM in a new mouse model of staphylococcal infection. The efficacy of lysostaphin was confirmed in the new model as well. Surprisingly, the catalytic domain of LytM was no more effective than control. This finding prompted us to compare properties of the two proteins in greater detail in vitro. Here, we report the in vivo observations and the in vitro properties of lysostaphin and LytM that might explain the different treatment outcomes.

**Results**

**Chronic contact eczema model of staphylococcal infection**

A new chronic dermatitis model of staphylococcal infection for in vivo functional studies was developed. Following standard procedures, mice were sensitized by epicutaneous application of 4-ethoxymethylene-2-phenyloxazolone (oxazolone, Sigma) on the abdomen skin. Six days later and subsequently every second day they were challenged with oxazolone applied to the ears. The treatment led to the development of chronic contact eczema in the treated ear, but not in the contralateral ear, which was left untreated as a control (Additional file 1).

Preliminary experiments were run to establish a suitable S. aureus dose for the infection experiments. 10$^6$, 10$^7$, 10$^8$, and 10$^9$ CFUs of S. aureus strain LS-1 were spread on both ears of one mouse each. Mice were sacrificed two days later, ears were homogenized and S. aureus colony forming units (CFUs) counted. 10$^6$ S. aureus cells per ear were sufficient to establish infection in oxazolone-treated, inflamed mouse ears, but not in non-oxazolone treated ears (data not shown).

To establish the time course for the infection, 10$^6$ S. aureus cells were applied to the oxazolone-treated, inflamed ears and to the non-oxazolone treated, contralateral control ears. At different time points following inoculation, mice were sacrificed, ears homogenized and S. aureus colony forming units (CFUs) counted. In non-oxazolone treated control ears, no bacteria were found after the application of 10$^6$ S. aureus cells. In oxazolone pretreated ears, colony counts peaked two days after exposure, and bacteria were almost fully cleared six days after the inoculation (Figure 2A).

**Lysostaphin is effective in the contact eczema model**

**LytM185-316 is not**

The newly developed eczema model was used for in vivo comparison of lysostaphin and LytM efficacies. 30 mice were divided into three groups of 10 mice each. All mice were sensitized to develop eczema, and subsequently had 10$^6$ CFUs of S. aureus P1 cells applied to their ears to induce dermatitis. Twelve hours after inoculation of bacteria the treatment with lysostaphin and LytM185-316 was started. 100 μg of lysostaphin or LytM185-316 in 50 mM glycine pH 8.0 with 10% glycerol was applied
topically to each mice ear in a volume of 20 μl. In the control group, buffer alone was used for the treatment. Ears were treated with proteins or buffer four times every 12 hours. Three hours after the last treatment mice were anesthetized, the ears dissected and the extent of infection estimated as described above. On average, the lysostaphin treatment reduced the colony count by roughly a factor of 10. In contrast to lysostaphin, LytM185-316 had no beneficial effect and was no better than control (Figure 2B). We reasoned that the different treatment outcomes could reflect differences in protein stability, affinity to either peptidoglycan or other components of cell walls, or the preference for a particular pH or ionic milieu and proceeded to test the influence of all these factors in vitro.

**Lysostaphin is proteolytically more stable than LytM185-316**

During treatment, lysostaphin and LytM185-316 were exposed both to bacterial proteases and to host proteases at the site of infection. Initial experiments demonstrated that both enzymes were stable in bacterial cultures (CFU ~10^6). The stability to host enzymes was tested with blood and serum from rat, which were available from unrelated experiments without the sacrifice of additional animals. After 4 h incubation in 5% blood, the majority of LytM185-316 was degraded while the degradation of lysostaphin was minimal. Both proteins were more stable in 5% serum, but again LytM185-316 was less stable than lysostaphin (Additional file 2).

**Lysostaphin and LytM185-316 recognize different cell wall components**

The affinity of lysostaphin and LytM was compared in a pulldown assay using various cell wall preparations that were increasingly enriched in peptidoglycan (Figure 3). Cell walls were used either crude (lane 2) or subjected to an extra washing step (lane 3), to SDS treatment, which should remove lipid components (lane 4), to TCA treatment, which is thought to remove teichoic acids (lane 5), or to trypsin treatment, which can be expected to remove protein components from cell walls (lane 6). The pulldown assay was also carried out with "purified" peptidoglycan, which was obtained from crude cell wall preparations by a combination of the SDS-, TCA- and trypsin treatments (lane 7), and with peptidoglycan from a commercial source (Fluka) (lane 8).

In all cases, lysostaphin bound to the cell wall preparations albeit with different efficiency. Our results suggest that binding to crude cell walls was most effective, probably because of interactions between lysostaphin and non-peptidoglycan components of *S. aureus* cell walls (Figure 3A).
In contrast, LytM\textsubscript{185-316} was not efficiently pulled down by crude cell wall preparations. However, when the cell walls were subjected to a washing step prior to the pulldown experiment, LytM\textsubscript{185-316} could be effectively pulled down. The effect of the washing step on the cell wall preparations is not clear. It may simply reduce clumping and make cell wall structures more accessible. Alternatively, it may remove a putative inhibitory factor in the unwashed cell wall sonicate. Further purification of peptidoglycan had a little effect on the outcome of the pulldown experiments. Therefore, we conclude that LytM\textsubscript{185-316} binds directly to cell walls and interacts primarily with peptidoglycans, rather than with other cell wall components (Figure 3B).

Full length LytM (without predicted signal peptide, LytM\textsubscript{26-316}) was not efficiently pulled down by any of the peptidoglycan preparations. Traces of protein were detected in the pulldown fraction in some cases, but the effect was probably unspecific, because no systematic trend with increasing peptidoglycan purity was observed (Figure 3C).

**Lysostaphin and LytM\textsubscript{185-316} bind peptidoglycan or cell walls differently**

The involvement of different regions of lysostaphin in peptidoglycan binding has been investigated earlier. The results show that lysostaphin has affinity for the pentaglycine crossbridges themselves [34], but also binds cell walls via the cell wall targeting domain [35]. In contrast, almost nothing is known about the role of different LytM fragments in peptidoglycan binding. Therefore, we investigated this question by the pulldown assay (Figure 4A). Comparing the amounts of protein in the pulldown and supernatant fractions, we found that the full length protein (LytM\textsubscript{26-316}) did not efficiently bind to peptidoglycan. Mutation of the Zn\textsuperscript{2+} ligand Asn117 to alanine, which should weaken the binding of the occluding region to the catalytic domain, did not significantly change the situation. The isolated N-terminal domain of the enzyme also failed to bind to peptidoglycan, whereas LytM\textsubscript{185-316} bound efficiently. When the two Zn\textsuperscript{2+} ligands His210 and Asp214 were separately mutated to alanine, the binding was lost again. Changing the third Zn\textsuperscript{2+} ligand, His293 of the HxH motif to alanine, made the protein insoluble as reported earlier [12], so that peptidoglycan binding could not be tested. The first histidine of the HxH motif, His291, is likely to act as a general base in catalysis [11]. When this residue was mutated to alanine, peptidoglycan binding was reduced, but not fully abolished.

The requirement of an intact active site for peptidoglycan binding was also supported by inhibitor studies. We had previously shown that EDTA and 1,10-phenanthroline blocked activity, presumably by chelating Zn\textsuperscript{2+} ions. We now observed that both metal chelators also abolished binding of LytM\textsubscript{185-316} to peptidoglycan (Figure 4B, lanes 1–2). In contrast, the weak Zn\textsuperscript{2+} ion chelator glycine hydroxamate and other small molecules and protease inhibitors did not interfere with peptidoglycan binding (Figure 4B, lanes 3–6). We conclude from these experiments that the accessibility and integrity of the active site is essential for the binding of the protein to peptidoglycan (Figure 4).

**Lysostaphin and LytM\textsubscript{185-316} activities depend differently on pH**

Peptidoglycan hydrolase activities were assayed in a turbidity clearance assay, using *S. aureus* cells. Perhaps due to remaining peptidoglycan hydrolase activity in the cell wall, there was some decrease of turbidity also in control, in the absence of exogenously added enzyme. Therefore, all apparent OD values at 595 nm were expressed as
percent of the control. A value close to 100% indicates a very low activity, whereas a very low OD reports highly active enzyme. Both lysostaphin and LytM185-316 were only marginally effective at pH 6.0 (50 mM phosphate buffer), but became much more active at pH 7.0. A further pH increase to the range between 7.0 and 9.0 (50 mM Tris–HCl) had little effect on the activity of lysostaphin, but enhanced the activity of LytM185-316. Even at pH 9.0, incubation with LytM185-316 lysed fewer cells than incubation with the equivalent amount of lysostaphin, particularly at late time points, possibly because of the lower stability of LytM185-316 (Figure 5).

**Lysostaphin and LytM185-316 activities depend very differently on ionic strength**

Investigating the pH dependence, we noticed a dramatic dependence of the lysis efficiency on the buffer. For example, the activity of LytM185-316 was much higher in 20 mM than in 50 mM Tris–HCl (both pH 8.0), and increased further when Tris was replaced with glycine at pH 8.0. However, glycine did not seem to act as an allosteric activator, because it did not enhance the activity when it was added in the presence of other buffer substances. Similar observations were made with other buffer components (Additional file 3).

A clear pattern emerged only when lysis activities of LytM185-316 and lysostaphin were correlated with the conductivity of the buffers (Figure 6). Lysostaphin degrades *S. aureus* cell walls inefficiently in low conductivity buffers, but becomes more efficient in buffers of higher conductivity. In contrast, LytM185-316 works best at low conductivity, and is almost ineffective in high conductivity buffers. The transition region for both effects is around 2 mS/cm, which corresponds roughly to a total ion concentration of 15–20 mM for singly charged cations and anions and typical mobilities (Figure 6).

Conductivity reflects both ion concentration and mobility. We reasoned that ionic strength was more likely than conductivity to influence protein activity, and therefore varied conductivity systematically by changing the concentration of sodium chloride between 0 and 500 mM. Lysostaphin and LytM185-316 activities were again dependent on the ionic strength in the expected manner, but conductivity was more directly correlated with ionic strength in this experiment (Figure 7).

The influence of ionic strength could also be demonstrated in a different way that was more directly related to the *in vivo* experiments. The low lytic efficiency of lysostaphin in glycine buffer could be overcome by addition of 25 to 100% of serum. Conversely, the addition of 25% or more serum to optimal reaction conditions for LytM185-316 (50 mM glycine-NaOH) completely abolished the activity of enzyme (data not shown).

The analysis of MIC and MBC for lysostaphin and LytM185-316 confirmed the above conclusions. The MIC for lysostaphin was around 0.0015–0.003 μg/ml, but inhibition of bacterial growth was not observed even with 5 μg/ml of LytM185-316. The MBC of lysostaphin was approximately 0.15 μg/ml in CASO broth and glycine buffer in agreement with previous data [36]. LytM185-316...
had an MBC around 0.3 μg/ml in the low ionic strength glycine buffer, but did not exhibit bactericidal activity in CAMH or CASO broth growth media which have conductivity 18 mS/cm.

Discussion
Lysostaphin treatment of *S. aureus* infection has been reported earlier. In a cotton rat model, *S. aureus* nasal colonization has been eradicated by this enzyme [26]. In the mouse, *S. aureus* systemic infections have been successfully treated [25] and biofilms have been effectively eliminated from a catheterized jugular vein [24]. The chronic dermatitis model of staphylococcal infection reported in this paper differs significantly from the earlier models and therefore represents an independent confirmation for the efficacy of lysostaphin. The lack of efficacy of the LytM185-316 treatment was initially surprising in light of previously observed comparable activity of lysostaphin and LytM185-316, though in experiments carried out in low salt buffers. As a result of this work, we now know that LytM185-316 differs from lysostaphin in several ways that could all explain the outcome of the mouse experiments.

Stability
LytM185-316 is a very stable protein in extracts of *Escherichia coli*, where the protein can be effectively produced recombinantly. The protein is also stable against staphylococcal proteases, just like lysostaphin. However, there are stability differences in serum and blood. This would obviously be relevant if lysostaphin or LytM were used systemically. As we are not sure to what extent the proteolytic stabilities in blood or serum reflect the situation in tissues with eczema, the influence of this factor on the overall treatment income is not clear though should not be neglected.

Binding
Both lysostaphin and LytM185-316 bind the pentaglycine crossbridges of *S. aureus* peptidoglycan. Both proteins recognize the crossbridges themselves, probably at least in part by interactions with the active site cleft. Lysostaphin has an extra cell wall targeting (CWT) domain which provides affinity. There is no counterpart in LytM (or LytM185-316), and therefore we originally expected that the N-terminal domain of the full length protein might play a similar role, especially in the light of the homology to SsaA. However, our experiments argue against this possibility, because full length LytM does not bind peptidoglycan.

Modular structure
LytM185-316 binds purified peptidoglycan the most effectively. The opposite is true for lysostaphin, which seems to recognize other cell wall components as well. It has previously been reported that deletion of the CWT domain in lysostaphin does not interfere with the endopeptidase activity of the enzyme, but abolishes its ability to distinguish between *S. aureus* and *S. staphylolyticus* [37]. As the peptidoglycans of the two bacterial species are identical [38], it suggests the recognition of non-cell wall components by CWT. Irrespective of which part of the lysostaphin protein provides the affinity to non-peptidoglycan cell walls, the ability of the protein to bind to crude cell walls is clearly helpful to lyse intact cells and seems to provide lysostaphin with an advantage as a protein drug. LytM is an autolysin, which is produced by the cell and delivered to the cell wall from “inside” while lysostaphin is a bacteriocin that approach target cells from the “outside”. In the treatment model, the approach of the peptidoglycan hydrolases to cell walls is necessarily from the outside, again favouring lysostaphin over any LytM fragment.

Ionic milieu
Perhaps the most crucial factor to explain the different treatment outcomes is the very different response of the two proteins to the ionic milieu. We do not know the precise ionic milieu of the contact eczema model of *S. aureus* infection, but suspect that it belongs to the high ionic strength regime, which would certainly apply for serum. If this is true, the ionic milieu in the mouse eczema could explain differences in treatment outcomes between lysostaphin preferring higher concentrations of salts for its activity and LytM being strongly inhibited in such environment.
Conclusions
Perhaps the strongest predictor of the mouse experiments is the biology of lysostaphin and LytM. As a bacteriocin, lysostaphin is evolved for the lysis of \textit{S. aureus} cell walls. In contrast, LytM as an autolysin should be evolved to have its activity under tight control. We expected this to apply for the full length enzyme, but hoped to bypass this step by the artificial activation that removes the N-terminal domain and the occluding region. Apparently, this does not suffice, because there are differences at several other levels which reflect the different \textit{in vivo} roles of lysostaphin and LytM. We conclude that the use of LytM\textsubscript{185-316} as an antibacterial agent is a more remote possibility than originally envisaged and that efforts to develop antibacterial peptidoglycan hydrolases should perhaps be concentrated on proteins that act as bacteriocins rather than autolysins.

Methods

Bacterial cultures

Bacteria were grown in CASO broth (Fluka) at 37°C with strong aeration from a 100-fold dilution of overnight cultures. Three strains of \textit{S. aureus} were used in the studies. The LS-1 is an arthritogenic strain originally isolated from swollen mouse joint [39]. The 8325–4 strain is a derivative of NCTC 8325, which has been cured of resident prophages and has low production of coagulase and surface adhesions [40]. The P1 strain was isolated from a rabbit inoculated with ATCC 25923 and cured of resident prophages and has low production of coagulase and surface adhesions [40]. The P1 strain was used to develop the eczema model, while the P1 strain was used for all \textit{in vitro} assays (pulldown, lysis, stability). The susceptibilities of the 8325–4 and P1 strains towards LytM and lysostaphin were comparable.

Proteins

A fragment of DNA corresponding to the LytM\textsubscript{24-105} protein was amplified by PCR from the previously described full length LytM clone [12] inserted into the pET15mod vector and called pET15modLytM\textsubscript{24-105}. The construct was then transformed into E. coli strain BL21(DE3). Protein expression was induced during the logarithmic growth phase of the bacteria (OD\textsubscript{600} of 0.8) by the addition of 1 mM IPTG and continued for 4 h at 25°C. The recombinant protein was purified by affinity chromatography on a Ni\textsuperscript{2+}-loaded column (Qiagen), followed by gel filtration on a Sephacryl S-200 column (Amersham Bioscience). LytM\textsubscript{26-316}, LytM\textsubscript{99-316}, LytM\textsubscript{185-316} and all point mutants were expressed and purified as previously described [12]. Lysostaphin (mature form) was purchased from Sigma and used without further purification.

Antibodies

Polyclonal antibodies against LytM\textsubscript{185-316} were raised in rabbit (Pineda Antibody Service, Berlin, Germany). Antibody purification was performed by affinity to LytM\textsubscript{185-316} protein coupled to CNBr-activated Sepharose 4B (Amersham Bioscience) according to the manufacturer’s instructions. After washing, antibodies were eluted with 100 mM glycine pH 2.7. The pH of the eluent was immediately neutralized by the addition of 1/10 volume of 2 M Tris–HCl pH 8.0. The concentration of the antibodies in the eluent was estimated based on the absorption at OD\textsubscript{280}.

Western blot hybridization

Proteins separated by SDS-PAGE were transferred onto ECL membrane (Amersham Bioscience) by semidry transfer and then incubated with 0.5 μg/ml purified antibodies against LytM\textsubscript{185-316} protein. Goat anti-rabbit peroxidase-conjugated secondary antibodies (Sigma) were detected using Western Blot Luminol Reagent (Santa Cruz Biotechnology).

LytM stability

Supernatants from 1 ml cultures of \textit{S. aureus} at late exponential phase were concentrated, mixed with 2 μg of LytM\textsubscript{26-316} and incubated overnight at 37°C. Proteins were separated on SDS-PAGE and used for Western blot hybridization. To assess the stability of lysostaphin and LytM\textsubscript{185-316} in buffer with addition of blood or serum (from rat) enzyme was mixed with 5% or 50% blood or serum in 50 mM glycine pH 8.0, and incubated at 37°C. Protein samples were collected after 1 and 4 h, separated by SDS-PAGE and used for Western blot hybridization.

Cell wall treatment

Late exponential phase cultures of \textit{S. aureus} grown in CASO Broth medium were harvested by centrifugation, resuspended in buffer A (20 mM Tris–HCl pH 7.5) and autoclaved for 20 min. Crude extract was obtained after sonicating the cells for 3 min. The accessory wall polymers were removed by the following methods. SDS treated walls were boiled in 4% SDS for 30 min. Trypsinized walls were prepared by 8 h trypsin digest (0.5 mg/ml) at 37°C. Trichloroacetic acid (TCA) treatment was done by 48 h incubation in 10% TCA at 4°C. After each of these treatments, cell walls were extensively washed in buffer A. Purified peptidoglycans were prepared as described previously [12] by combining all methods described above. Alternatively, \textit{S. aureus} peptidoglycan was purchased from Fluka Biochemika.
Pulldown peptidoglycan binding assay
To assess binding, 2 μg of protein was mixed with cell walls or peptidoglycans (100 μg) and incubated at room temperature for 15 min. Then, soluble and insoluble fractions were separated by centrifugation and peptidoglycans were washed with 1 ml of buffer A. Soluble fractions and washed peptidoglycans were mixed with loading buffer separated by SDS-PAGE and analyzed by Western blot hybridization. Final concentrations of 10 mM EDTA, 1 mM 1,10-phenanthroline, 10 mM N-acetylglucosamine, 10 mM glycine hydroxamate, 1 mM PMSF and 1 mM E-64 were used to test the influence of these compounds on peptidoglycan binding.

Cell lysis assay
S. aureus cells collected at the exponential growth phase were washed and suspended in buffer A supplemented with 200 μg/ml erythromycin. Then the cells were diluted to an apparent OD595 of 1.8 with an appropriate buffer. Enzymes were added to the final concentration of 18nM and 200 μl of reaction transferred onto the microtiter plate. Plates were incubated at 37°C with 2 s shaking every 5 min. OD of the suspension was checked at the wavelength of 595 nm at 0, 20, 40, 60, 90 or 120 min. after starting the reaction. Ionic strength of the reaction milieu (cells resuspended in appropriate buffer) was measured using conductivity meter MeteLab CDM230 (Radiometer Analytical, France) at the beginning of the tests. Lytic activity was calculated as a per cent of control OD595 (the same samples as for reaction but without enzymes). Each experiment was repeated twice in quadruplicate.

Determination of minimal inhibitory concentration (MIC) and minimal bactericidal concentration (MBC)
Both parameters were determined generally as described by Kusuma and Kokai-Kun [36]. For MIC determination by the microdilution method, 100 μl of Cation-Adjusted Mueller-Hinton broth were inoculated with ~104 S. aureus cells (strain 8325–4) and enzyme concentrations between 4 and 0.0015 μg/ml were tested. For MBC determination, ~106 CFU/ml of S. aureus cells (strain 8325–4) in either CASO broth or in 50 mM glycine pH 7.5 were incubated with between 10 to 0.15 μg/ml of enzyme. For lysostaphin, but not for LytM, the buffer was supplemented with 150 mM NaCl to make digestion conditions optimal for the enzyme.

Animal experiments
Ethical permission for animal experiments was obtained from the Animal Research Ethics Committee of Göteborg University. Throughout the experiments the animals were under control of the veterinarian. No differences in animal behavior and general state of health were observed between the control and experimental groups.

Induction of chronic contact eczema in mice
NMRI mice were sensitized by epicutaneous application of 150 μl of a mixture of ethanol and acetone (3:1) containing 3% of 4-ethoxymethylene-2-phenyloxazolone (oxazolone, Sigma) on the abdomen skin. Six days later, and subsequently every second day, all the mice were challenged on both sides of one ear with 30 μl 1% oxazolone dissolved in olive oil. The mice received altogether 4 oxazolone challenges on the ear. This procedure leads to chronic, eczematous skin inflammation characterized macroscopically by swelling, redness and superficial desquamation and microscopically by influx of inflammatory cells (Additional file 4).

Infection kinetics
The day following the last application of oxazolone, the mice were briefly anaesthetized, and S. aureus in a volume of 10 μl was spread on the skin surface of the inflamed ear. In the first experiment four mice with dermatitis were subjected to skin infection in one ear while the contra lateral ear was used as a control. S. aureus strain LS-1 at 106, 107, 108, and 109 CFU (colony forming unit) was spread on each ear, and the mice were sacrificed two days later. In the second experiment, the kinetic of infection was assessed. Twenty mice with dermatitis on one ear were exposed to 106 CFU S. aureus strain LS-1. Groups of five mice each were sacrificed at 1, 2, 3, and 6 days following exposure to bacteria. One mouse ear/group was subjected to histological examination (Additional file 4) and the rest 4 ears/group were subjected to enumeration of staphylococci.

Comparison of lysostaphin and LytM185-316 in the mouse model
In the last in vivo experiment the staphylococcal strain P1 (106/ear) was used to infect ears of mice with eczema. Twelve hours after inoculation of bacteria the treatment with proteins was started; 100 μg of lysostaphin or LytM185-316 in 50 mM glycine pH 8.0 and 10% glycerol buffer was applied to each mouse ear in a volume of 20 μl. In the case of control mice buffer alone was used for the treatment. Ears were treated with proteins or buffer four times every 12 hours. Three hours after the last treatment mice were anesthetized and the ears dissected. The ears were washed with alcohol to remove surface bound bacteria, kept on ice, homogenized and diluted in PBS. One hundred microliter of the homogenate from various dilutions was then transferred to agar plates, containing 7.5% sodium chloride. After incubation at 37°C for 24 hours the colony forming units were counted. 10 mice were used in the control group and in each treatment group.

Prior to the in vivo use, staphylococci were cultured for 24 hours on blood agar plates, re-inoculated and
grown on fresh blood agar plates for another 24 hours, harvested, and stored frozen at −20°C after suspending aliquots in phosphate-buffered saline (PBS) supplemented with 5% bovine serum albumin and 10% dimethyl sulfoxide. Before application on ears, staphylococcal suspensions were thawed, bacteria washed in PBS and diluted in PBS to achieve the appropriate concentration of the staphylococci. To determine the CFU, aliquots of staphylococcal suspensions were subjected to dilution, plating on blood agar and enumeration.

Acknowledgements

We are thankful to Drs Renata Filipek and Ezbicka Nowak for critical reading of the manuscript and fruitful discussions. This work was supported by the European Communities ("Novel non-antibiotic treatment of staphylococcal diseases", specific RTD program QLRT-2001-01250, Center of Excellence in Bio-Medicine, EC FP7 grant "Proteins in Health and Disease" (HEALTH-PROT, 2008, 517(7):1981–1990). 2008, 517(7):1981–1990).

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Additional file

Additional file 1: Picture of mouse ears untreated (on the left) and treated (on the right) with oxazolalone.

Additional file 2: Stability of LytM155–316 and lysostaphin. Proteins were incubated without (1) or with concentrated, conditioned S. aureus media (2), 5% (3) or 50% (5), blood and 5% (6) or 50% (7) serum. After incubation proteins were separated by SDS-PAGE electrophoresis and detected by Western blot hybridization with anti-LytM antibodies.

Additional file 3: Time course of S. aureus 8325–4 cell lysis by LytM155–316 and lysostaphin in various conditions. (A) Influence of glycine. Lysis experiments were done in 100 mM glycine-NaOH, pH 8.0, 50 mM Tris-HCl, pH 8.0 and 100 mM glycine in 50 mM Tris-HCl pH 8.0. (B) Influence of mono-, di- and triglycine. Buffers were made as 50 mM with pH adjusted to 8.0 with NaOH. For comparison lysis in dd water was also checked. (C) Influence of various aminocoids. 50 mM L-arginine-HCl, DL-alanine-NaOH, L-arginine-HCl, L-glutamic acid-NaOH, diaminopimelic acid (DAP)-NaOH of pH 8.0 were tested. Lysis experiments were performed as described in Material and Methods.

Additional file 4: Histological examination of mouse ear during the development of eczema and S. aureus infection. (A) section of control ear, (B) section 2 days after S. aureus infection; massive invasion of inflammatory cells can be observed (indicated with open arrows).
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