Functional role of surface layer proteins of *Lactobacillus acidophilus* L-92 in stress tolerance and binding to host cell proteins

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*Lactobacillus acidophilus* surface layer proteins (SLPs) self-assemble into a monolayer that is non-covalently bound to the outer surface of the cells. There they are in direct contact with the environment, environmental stressors and gut components of the host in which the organism resides. The role of *L. acidophilus* SLPs is not entirely understood, although SLPs seem to be essential for bacterial growth. We constructed three *L. acidophilus* L-92 strains, each expressing a mutant of the most abundant SLP, SlpA. Each carried a 12-amino acid c-myc epitope substitution at a different position in the protein. A strain was also obtained that expressed the SlpA paralog SlpB from an originally silent *slpB* gene. All four strains behaved differently with respect to growth under various stress conditions, such as the presence of salt, ox gall or ethanol, suggesting that SlpA affects stress tolerance in *L. acidophilus* L-92. Also, the four mutants showed differential in vitro binding ability to human host cell proteins such as uromodulin or dendritic cell (DC)-specific intercellular adhesion molecule-3 grabbing non-integrin (DC-SIGN). Furthermore, co-culture of murine immature DCs with a mutant strain expressing one of the recombinant SlpA proteins changed the concentrations of the cytokines IL-10 and IL-12. Our data suggest that SlpA and SlpB of *L. acidophilus* participate in bacterial stress tolerance and binding to uromodulin or DC-SIGN, possibly leading to effective immune-modification.

Key words: *Lactobacillus acidophilus*, surface layer proteins, stress tolerance, probiotics, immune regulation

INTRODUCTION

*Lactobacillus acidophilus* is one of the most studied and consumed probiotic bacteria because of its high ability to survive in digestive juice, to bind to gut epithelial cells and to modulate host immune function [1, 2]. Our group has studied the effect of *L. acidophilus* L-92 mainly on immune function. Oral administration of *L. acidophilus* L-92 [3] has been reported to improve symptoms of atopic dermatitis [4–6], pollen allergy [7] and perennial allergic rhinitis [8] in humans and mice. In addition, oral intake of the strain protected against influenza infection in the mouse and human [9, 10]. Strain L-92 has immune-regulating activities in mice, such as the regulation of cytokine production, suppression of antigen-specific IgE, induction of apoptosis of antigen-stimulated T cells and induction of regulatory T cells [11–13].

*L. acidophilus* produces surface layer proteins (SLPs) that form a self-assembled monolayer on the outer surface of the cells [14]. Given this location, these proteins might be important for contact with the environment [15]. *L. acidophilus* can potentially produce 3 SLPs, SlpA, SlpB and SlpX, in addition to expressing various surface layer-associated proteins (SLAPs) [16]. SlpA is the most abundant SLP, as SlpX is only expressed at low amounts and SlpB is not expressed because the gene does not have a functional promoter [17]. An *L. acidophilus* NCFM slpX knockout strain showed both a lower survival rate under specific growth conditions and lower binding to mucin [18]. These observations are indicative of the importance of (certain) SLPs for environmental stress tolerance and the binding to components of host cells. An *L. acidophilus* NCFM mutant carrying an *slpA* knockout insertion expressed SlpB, which was caused by
an inversion between the originally silent slpB gene and slpA, resulting in slpB being under the control of the slpA promoter in this strain [19, 20]. These results indicate that L. acidophilus cells need to express at least SlpA or SlpB on their surface in order to grow and survive and that SLPs might be very important or even essential in L. acidophilus.

SlpA is one of the key components in L. acidophilus binding to host proteins and further immune-modulation. L. acidophilus CP23, which has a relatively low level of SLPs on its surface, shows lower adhesion activity to caco-2 cells and less IL-12 release from DCs than L. acidophilus strains possessing more SLPs [21, 22]. Immune regulation by L. acidophilus NCFM SlpA operates via its binding to one of the C-type lectin receptors, the specific intracellular adhesion molecule-3 grabbing non-integrin homolog-related 3 (SIGNR3) in the mouse and dendritic cell (DC)-specific intercellular adhesion molecule-3 grabbing non-integrin (DC-SIGN) in humans [19, 23]. In addition, the uromodulin expressed in microfold (M) cells selectively binds to SlpA on L. acidophilus L-92 cells [24]. Binding leads to selective incorporation of the SlpA-expressing bacteria into the M cells and to their effective delivery to DCs.

In addition to SLPs, several SLAPs have been shown to affect the organism’s binding activity to host components and immune-modulating activity [16, 25, 26]. Deletion of PrtX (encoded by LBA1578) was reported to increase adhesion of the L. acidophilus NCFM mutant to mucin and fibronectin [25], while on the other hand, deletion of another SLAP, LBA0191, decreased the adhesion to both human proteins [26]. When murine DCs were exposed to cells lacking PrtX, the concentrations of the pro-inflammatory interleukins IL-6 and IL-12, that of the anti-inflammatory interleukin IL-10, and the IL-10/IL-12 ratio were increased compared with those of the parent strain. Increase of inflammatory interleukin IL-12, and the IL-10/IL-12 ratio were decreased while on the other hand, deletion of another SLAP, LBA0191, decreased the adhesion to both human proteins [26]. When murine DCs were exposed to cells lacking PrtX, the concentrations of the pro-inflammatory interleukins IL-6 and IL-12, that of the anti-inflammatory interleukin IL-10, and the IL-10/IL-12 ratio were increased compared with those of the parent strain. Increase of the IL-10/IL-12 ratio suggests a pro-inflammatory status. The LBA1029 deletion mutant caused lower induction of TNF-α in murine DCs than its parent [16].

In the present study, we constructed four genetically modified L. acidophilus L-92 SlpA mutants by inserting a c-myc epitope at different positions in the protein and investigated the roles of certain regions in SlpA on both bacterial stress tolerance and binding to host proteins such as uromodulin and DC-SIGN.

**MATERIALS AND METHODS**

**Bacterial strains and growth conditions**

The L. acidophilus L-92 used was from the stock collection of Asahi Group Holdings Ltd. L. acidophilus strains were grown in de Man, Rogosa and Sharpe (MRS) medium (BD, Franklin Lakes, NJ, USA) at 37°C. In general, 0.5 to 5% of a culture was inoculated in fresh medium and incubated overnight at 37°C without shaking. Lactococcus lactis and Escherichia coli were used for cloning purposes. L. lactis was cultured at 30°C in Difco M17 medium (BD) containing 0.5% glucose. E. coli was cultivated in Luria Bertani (LB) medium (Formedium, Norfolk, UK) at 37°C with shaking. The media were supplemented with 5 µg/mL chloramphenicol (Cm; for Lactobacillus and Lactococcus) or 10 µg/mL Cm (for E. coli), and when needed, they were further supplemented with 5 µg/mL erythromycin (Ery; for Lactobacillus and Lactococcus) or 150 µg/mL Cm (for E. coli). Strains, plasmids and primers used in this study are listed in Supplementary Tables 1 and 2. Cloning, transformation and mutagenesis strategies are described in detail in the Supplementary Materials and Methods online.

**SDS-PAGE and Western blotting**

Tricin sodium dodecyl sulfate-10% polyacrylamide gel electrophoresis (SDS-PAGE) and Coomassie brilliant blue (CBB) staining were performed as described previously [27]. Cells cultured overnight at 37°C in MRS (with 1 µg/mL Ery only for SA2r) were washed and resuspended in PBS to adjust the OD<sub>600</sub> to 4.8. Cell solutions (30 µL) were mixed with 10 µL of Laemmli sample buffer (Bio-Rad Laboratories) and then boiled for 10 min, after which 10 µL of the samples were applied to the SDS-polyacrylamide gel. Proteins were blotted onto a polyvinylidene fluoride (PVDF) membrane (Bio-Rad Laboratories), after which immune detection of recombinant SlpA containing the c-myc epitope was carried out with an anti-c-myc mouse monoclonal antibody (Thermo Fisher Scientific), horseradish peroxidase (HRP)-conjugated anti-mouse IgG antibody (Thermo Fisher Scientific) and chemiluminescent detection kit (ECL Prime kit, Bio-Rad Laboratories) according to manufacturers’ manuals. Samples from some protein bands were purified from gel fragments and analyzed by nano LC/MS/MS (Applied Biosystems) at JPROteomics (Miyagi, Japan), as described previously [28]. The Mascot software was used to perform mass data acquisition.

**SLP and SLAP isolation**

SLPs and SLAPs were isolated essentially according to a previous report [16]. Cells grown overnight in MRS were harvested by centrifugation at 8,000 g for 5 min and washed twice with PBS. All subsequent steps were conducted at 4°C. Cells were resuspended in a 1/20 volume of 5 M LiCl, gently agitated for 15 min, and then centrifuged at 20,000 g for 30 min. The supernatants were dialyzed in a 10,000-Da membrane overnight against water. After dialysis, the solutions were centrifuged at 20,000 g for 30 min. The precipitates were then resuspended in 1 M LiCl and incubated at 4°C for 15 min, after which the suspensions were centrifuged at 20,000 g for 10 min. For SLP fractions, the precipitates were then washed three times with cold water and resuspended in water or 10% SDS. For SLAP fractions, the supernatants after centrifugation in the 1 M LiCl incubation step were dialyzed, centrifuged, washed and resuspended in water or 10% glycerol.

**Fluorescence microscopy and flow cytometry**

The c-myc epitope was detected with antibodies, fluorescence microscopy and flow cytometry basically as described previously [29, 30]. Cells were grown overnight in MRS at 37°C, harvested and washed with PBS. The washed cells were resuspended in PBS containing 0.25% bovine serum albumin (BSA) and the primary antibody (50-fold diluted mouse anti-c-myc antibody) and incubated on ice for 30 min. The cells were then washed twice with PBS and resuspended in PBS with 0.25% BSA and the 100-fold diluted secondary antibody (goat anti-mouse IgG secondary antibody, FITC, Thermo Fisher Scientific) and incubated on ice for 30 min. Subsequently, the cells were washed with PBS and analyzed by fluorescence microscopy and flow cytometry. Fluorescent levels in approximately 50,000 cells were measured with a BD FACSCanto flow cytometer (BD Biosciences, Franklin Lakes, NJ, USA) using a 488-nm argon laser. Raw data were obtained using the provided software and analyzed using FlowJo (BD Biosciences).
collected using the FACS DIVA software (BD Biosciences). WinMDI 2.9 was used for data analysis (http://en.bio-soft.net/other/WinMDI.html). Microscopy images were taken with a DeltaVision microscope (Applied Precision, Issaquah, WA, USA) equipped with a 2300-W xenon light source, 100× bright field objective, and a GFP filter set (Chroma, Bellows Falls, VT, USA; excitation at 470/40 nm and emission at 525/50 nm).

**Determination of growth and stress tolerance**

*L. acidophilus* strains were cultured overnight in MRS medium. The overnight culture of each strain was diluted in fresh MRS medium with or without 0.005% SDS, 0.3% ox gall, 5% ethanol (EtOH) or 2% sodium chloride (NaCl) to an OD_{600} of around 0.03 and incubated at 37°C in the wells of a 96-well microtiter plate with a transparent lid in a Powerscan HT microplate reader (DS Pharma Biomedical, Osaka, Japan). The OD_{600} was measured every hour after 10 sec of shaking.

**DC-SIGN and uromodulin binding assay**

*L. acidophilus* cells grown in MRS for 20 hr were harvested and washed with the same volume of PBS twice and then resuspended in the same volume of 0.1% trisodium citrate. The cell suspension was heated at 95°C for 10 min to kill the cells. Subsequently, they were washed with PBS and resuspended in PBS while adjusting the cell concentration to the desired density for the binding and co-culturing assays. The binding ability of *L. acidophilus* strains to DC-SIGN and uromodulin was analyzed using ELISA as described previously [19]. A recombinant human DC-SIGN/CD209 Fc chimera protein (R&D Systems, Minneapolis, MN, USA) and recombinant mouse uromodulin Fc chimera protein [24] were used. Fifty microliters of the heat-killed bacterial cells (7.5 × 10^9 cells/mL) were coated on NUNC 96-well MaxiSorp plates (Thermo Fisher Scientific) and incubated overnight at 4°C. The plates were blocked with 1% of BSA in PBS for 1.5 hr. DC-SIGN (5 μg/mL) or uromodulin (5 μg/mL) in PBS was added, and the plates were then incubated for 1 hr with gentle shaking. Bound protein was detected with an HRP-conjugated goat anti-human IgG-Fc fragment antibody (Bethyl Laboratories, Montgomery, TX, USA) and TMB One Component HRP (Montgomery, TX, USA) and the presence and identity of the recombinant *L. acidophilus* gene was confirmed using the FACSDiva software (BD Biosciences). At least 10^6 events per condition were acquired. The statistical significance of differences between the parent strain and each mutant was calculated using 2-way ANOVA in SPSS and the post hoc paired t-test with Bonferroni correction.

**RESULTS**

**Construction of *L. acidophilus* L-92 expressing recombinant SlpA**

As the *slpA* gene seems to be essential for *L. acidophilus* L-92 and cannot be deleted (data not shown), we decided to make mutant strains expressing recombinant forms of SlpA. A stretch of ten amino acid residues in non-conserved regions in SlpA was replaced with 12 amino acid residues containing the c-myc epitope as an expression tag. The recombinant genes were integrated at the *slpA* locus using double crossover gene replacement recombination (DCO; Table 1). From a total of 10 different recombinant *slpA* genes that we attempted to make, only 3 could be obtained that were expressed (Table 1). Two mutant proteins, SlpAr2 and SlpAr3, carried the c-myc epitope in an area in the SlpA protein that is predicted to be surface exposed [29, 32]. The other two, SlpAr1 and SlpAr4, carried the substitution in a structurally unspecified region. In total, four mutants were obtained in *L. acidophilus* L-92 Δupp, a strain in which the *upp* gene was deleted to enable using the *upp*-based counterselection system [18]. DCO, as confirmed by PCR results showing replacement of the original *slpA* gene by the recombinant copy of *slpA*, was attained in three cases (strains SAr1, SAr4 and SAr3; SAr3 renamed to SB3; see below). In one strain, designated SAr2, only single crossover (SCO) could be obtained such that both the original *slpA* gene and the recombinant copy, *slpAr2*, were present in the chromosome. The SAr2 strain was used for further studies, as excision of the integrated plasmid, required for exchange of the *slpA* genes, was repeatedly unsuccessful. The *slpA* regions in the chromosomes of the three DCO mutants were sequenced, and the presence and identity of the recombinant *slpA* version was confirmed in two (SAr1 and SAr4). However, in one (SAr3, designated SB3 hereafter), the *slpB* gene was present at the *slpA* locus, while the recombinant *slpAr3* gene was at the silent *slpB* locus, suggesting that homologous recombination had occurred between the adjacent *slpA* and *slpB* genes. *L. acidophilus* L-92 *slpB* is a paralog of *slpA*; it is located in the opposite orientation around 3 kbp downstream of *slpA* (Supplementary Fig. 1). The *slpB* gene has two regions of identity with *slpA*. One area of around 300 bp is located in the 5'-UTR and the start of the coding regions; the other (approximately 400 bp) is present at the end of the 3'-UTR.

**Incubation of dendritic cells with recombinant *L. acidophilus* strains**

Animal experiments were approved by the Institutional Animal Experiment Committee (Application No. 16-28-01) of Asahi Group Holdings, Ltd. on December 2, 2016, and performed in 2017 at the animal facility of Asahi Group Holdings, Ltd. Female BALB/c mice (5 weeks old) were purchased from Japan SLC. Murine immature dendritic cells were prepared from bone marrow cells as described previously [31]. Bone marrow from mice femurs was flushed using cold PBS. Cells were treated with Tris-buffered ammonium chloride to lyse erythrocytes. The remaining cells were cultured in RPMI 1640 complete medium plus 10% fetal bovine serum (FBS) with mouse granulocyte-macrophage colony-stimulating factor (GM-CSF; 20 ng/mL) in NUNC six-well tissue culture plates (Thermo Fisher Scientific).

After 6 hr of incubation, non-adherent cells were removed, and the remaining cells were cultured in fresh RPMI 1640 complete medium plus 10% FBS containing GM-CSF for 7 days. Every other day, cultures were fed the fresh medium containing GM-CSF. On day 7, cells were harvested and, after confirmation that more than 90% were CD11c positive, they were treated with heat-killed *L. acidophilus* cells at a DC-to-*L. acidophilus* ratio of 1:10 or 1:1. After 24 hr of incubation, the supernatants were sampled to measure cytokines with a Bio-Plex Pro Mouse Th1 7-plex kit (Bio-Rad Laboratories) according to the manufacturer’s protocol. DCs (5 × 10^5) were incubated with surface marker monoclonal antibodies for 30 min at 4°C, washed extensively with BD Stain Buffer (BD Biosciences) and analyzed using a FACSCanto flow cytometer (BD Biosciences). At least 10^6 gated events were acquired. The statistical significance of differences between the parent strain and each mutant was calculated using 2-way ANOVA in SPSS and the post hoc paired t-test with Bonferroni correction.

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As the *slpA* gene seems to be essential for *L. acidophilus* L-92 and cannot be deleted (data not shown), we decided to make mutant strains expressing recombinant forms of SlpA. A stretch of ten amino acid residues in non-conserved regions in SlpA was replaced with 12 amino acid residues containing the c-myc epitope as an expression tag. The recombinant genes were integrated at the *slpA* locus using double crossover gene replacement recombination (DCO; Table 1). From a total of 10 different recombinant slpA genes that we attempted to make, only 3 could be obtained that were expressed (Table 1). Two mutant proteins, SlpAr2 and SlpAr3, carried the c-myc epitope in an area in the SlpA protein that is predicted to be surface exposed [29, 32]. The other two, SlpAr1 and SlpAr4, carried the substitution in a structurally unspecified region. In total, four mutants were obtained in *L. acidophilus* L-92 Δupp, a strain in which the *upp* gene was deleted to enable using the *upp*-based counterselection system [18]. DCO, as confirmed by PCR results showing replacement of the original *slpA* gene by the recombinant copy of *slpA*, was attained in three cases (strains SAr1, SAr4 and SAr3; SAr3 renamed to SB3; see below). In one strain, designated SAr2, only single crossover (SCO) could be obtained such that both the original *slpA* gene and the recombinant copy, *slpAr2*, were present in the chromosome. The SAr2 strain was used for further studies, as excision of the integrated plasmid, required for exchange of the *slpA* genes, was repeatedly unsuccessful. The *slpA* regions in the chromosomes of the three DCO mutants were sequenced, and the presence and identity of the recombinant *slpA* version was confirmed in two (SAr1 and SAr4). However, in one (SAr3, designated SB3 hereafter), the *slpB* gene was present at the *slpA* locus, while the recombinant *slpAr3* gene was at the silent *slpB* locus, suggesting that homologous recombination had occurred between the adjacent *slpA* and *slpB* genes. *L. acidophilus* L-92 *slpB* is a paralog of *slpA*; it is located in the opposite orientation around 3 kbp downstream of *slpA* (Supplementary Fig. 1). The *slpB* gene has two regions of identity with *slpA*. One area of around 300 bp is located in the 5'-UTR and the start of the coding regions; the other (approximately 400 bp) is present at the end of the 3'-UTR.
of the coding regions of both genes. As a result of the presumed homologous recombination event, the entire \textit{slpB} gene is under the control of the \textit{slpA} promoter in strain SB3. The \textit{slpAr3} gene is not expressed, as the \textit{slpB} locus does not carry an active promoter \[33\]. The four mutants were all used in further experiments.

Recombinant SlpA proteins are expressed in \textit{L. acidophilus}

Washed whole-cell suspensions of all four recombinant strains and their parent were boiled and subjected to SDS-PAGE, after which the gels were blotted and examined by Western analysis using anti-c-myc antibodies. As expected, all strains produced proteins in the size range of SlpA of approximately 43 kDa (Fig. 1). The recombinant and original SlpA proteins could not be distinguished on the gel, as they had almost the same sizes. Strain SB3 also produced such a protein, but it did not react with the c-myc antibody. This protein was a surface layer protein that was abundant on SB3 cells, and it was identified as SlpB through peptide mass fingerprinting (see below). Furthermore, SCO strain SAr2 showed a weak antibody response, suggesting that the amount of recombinant SlpA-c-myc protein (SlpAr2) produced by this strain was less than that made by SAr1 and SAr4. Most likely, strain SAr2 produces a mixture of SlpA and recombinant SlpAr2. The DCO strains SAr1 and SAr4 express their respective recombinant SlpA proteins of the expected sizes, each of which reacts with the c-myc antibody.

The c-myc epitopes in recombinant SlpA are surface exposed

To examine whether the parts in SlpA carrying the c-myc epitope substitutions are at the outside of the cells and could thus be in contact with host cell components, their accessibility for the c-myc antibody in whole cells was investigated. Cells of the three recombinant SlpA-expressing strains were first coated with a mouse anti-c-myc antibody and then with a FITC-labeled anti-mouse IgG antibody. The strains SAr1 and SAr2 reacted with the antibodies and were detectable by both fluorescence microscopy and flow cytometry (Fig. 2). In other words, the c-myc epitopes in the surface layer formed by these recombinant SlpA proteins are accessible from the outside. Strain SAr4, like its parent \textit{L. acidophilus} L-92 \textit{Δupp}, did not react with the antibodies; apparently, the c-myc epitope is shielded in this strain.
Surface layer-associated proteins

Surface layer-associated proteins have previously been identified on *L. acidophilus* NCFM cells. They have also been shown to affect the organism’s binding activity to host components and immune-modulating activity [16, 25, 26]. As SLAPs are thought to be associated with SLPs [16], the amino acid changes in the recombinant SlpAs might affect the SLAP patterns of the *L. acidophilus* mutants studied here. The SLPs and SLAPs of the recombinant strains were isolated by LiCl extraction and analyzed by SDS-PAGE. The SLP fractions of *L. acidophilus* L-92 Δupp and three of the recombinant strains showed a single intense band of a protein of the size of SlpA, while two bands were present in the extract of strain SB3 (Fig. 3). The proteins in the latter two bands were isolated from the SDS-polyacrylamide gel and identified with peptide mass fingerprinting. The larger band was identified as SlpX, while the protein in the smaller band, running to the same position as SlpA in the other three samples, was SlpB. The precipitated proteins after dialysis of 5 M LiCl extract were washed with 1 M LiCl. The resulting precipitated proteins and soluble proteins in the 1 M LiCl washing solution were SLP and SLAP fractions, respectively. M, protein marker (the same in both gels). The sizes of protein markers are indicated in the left margin. Description of the strains/samples (Table 1).
Growth and stress tolerance of *L. acidophilus* L-92 expressing mutant SlpA

When the growth of *L. acidophilus* L-92 Δupp and its mutants was examined in the nutritionally rich MRS medium, no differences were observed between the growth rates of the strains (Fig. 4 and Table 2). It has been suggested that SLPs have a role in tolerance to environmental stressors such as detergent, salt and alcohol. Indeed, each mutant showed strain-specific stress tolerance patterns when exposed to various stress conditions. The lag time of SAr1 in MRS with NaCl was longer than that of the parent, while growth was comparable in other media (data not shown). The growth rate of SAr2 was greatly decreased in ox gall-containing MRS broth, and the final optical density was lower than that of the parent. The lag time of SB3, the strain expressing SlpB, was shorter than that of its parent under all stress conditions tested. Strain SAr4 grew slower in ethanol-containing medium than the parent.

**Binding of DC-SIGN and uromodulin to *L. acidophilus* L-92 and its mutants**

Next, ELISA was performed to evaluate the interaction of the three recombinant SlpA-expressing strains and the strain expressing SlpB and SlpX with two specific host proteins: mouse uromodulin Fc chimera protein and recombinant human DC-SIGN/CD209 Fc chimera protein (Fig. 5). The binding of DC-SIGN to heat-killed SAr1 and SAr4 cells was slightly decreased

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Table 2. Growth characteristics of *L. acidophilus* strains

|μMax (h⁻¹) | Aupp | SAr1 | SAr2 | SB3 | SAr4 |
|---|---|---|---|---|---|
| MRS | 0.65 | 0.56 | 0.56 | 0.68 | 0.51 |
| + EtOH | 0.41 | 0.36 | 0.35 | 0.44 | 0.23 |
| + NaCl | 0.45 | 0.38 | 0.41 | 0.51 | 0.43 |
| + Oxgal | 0.62 | 0.6 | 0.4 | 0.57 | 0.54 |
| + SDS | 0.51 | 0.54 | 0.43 | 0.62 | 0.5 |

|Lag time (hr) | Aupp | SAr1 | SAr2 | SB3 | SAr4 |
|---|---|---|---|---|---|
| MRS | 3.4 | 3.7 | 2.7 | 3.2 | 3.2 |
| + EtOH | 4.8 | 4.5 | 3.7 | 3.4 | 4.6 |
| + NaCl | 7.3 | 8.8 | 6.3 | 5.1 | 6.7 |
| + Oxgal | 4.4 | 4.3 | 2.9 | 2.5 | 4.2 |
| + SDS | 3.3 | 3 | 2.1 | 2.3 | 3.1 |

The data are extracted from the curves in Fig. 3.
relative to the parent. Compared with the parent strain, binding of uromodulin to all three recombinant SlpA-producing mutants was significantly decreased, while it bound better to strain SB3. This result suggests that expression of SlpB and SlpX in strain SB3 does not affect the binding of DC-SIGN but instead strengthens the binding of uromodulin. The strains SAr1 and SAr4 showed decreased binding of both host proteins. The two strains express a recombinant SlpA variant in which the c-myc epitope is predicted to be, respectively, either surface exposed or not. Apparently, changes in the identity of SLPs as well as the structure of one of them, SlpA, can all affect binding of host proteins such as DC-SIGN and uromodulin.

Cytokine production from DCs is changed by L. acidophilus L-92 SAr2

Heat-killed recombinant L. acidophilus L-92 cells were incubated with immature dendritic cells (iDCs) derived from murine bone marrow to examine their role in cytokine production and maturation. Expression of three maturation markers, CD80, CD86 and MHCII, in iDCs after 24 hours of treatment with each of the mutant strains was not changed compared with that after treatment with the parent (data not shown), suggesting that they do not affect DC maturation. The concentration of 6 cytokines measured was not changed when iDCs were incubated with strain SAr1, SB3 or SAr4 compared with when they were incubated with the parent (Fig. 6). However, when the iDCs were incubated with strain SAr2, the concentrations of IL-10 and IL-12 were decreased compared with when the iDCs were incubated with the parent. These results may suggest that an alteration of SLPS and/or SLAPs on the mutant strain SAr2 changed DC signaling.

DISCUSSION

The four mutant L. acidophilus strains constructed here either express a recombinant version of the SlpA protein or a combination of SlpB and SlpX. SlpB is normally not expressed, as it is encoded by a silent gene without a functional promoter. The strains were used to demonstrate the importance of SLPs, of which at least one should be expressed on the surface of L. acidophilus, in bacterial stress tolerance and host protein binding, two critical parameters in the modulation of the host immune system.

SlpB is paralogous to SlpA, with 58% amino acid identity (Supplementary Fig. 2). Expression of SlpB (and SlpX, which is overexpressed in strain SB3) might allow the bacteria to grow better than when they would only express slpAr3, or the latter could even be lethal. Expression of small amounts of SlpX, next to SlpA, was detected in L. acidophilus strains L-92 (unpublished data) and NCFM [16], suggesting that all three recombinant strains obtained here might also express some SlpX but that the amount could not be determined by SDS-PAGE. Abundant expression of SlpX in the presence of SlpB, observed here for L. acidophilus SB3, was also reported in L. acidophilus ΔslpA expressing SlpB [18], indicating that SlpB cannot substitute for SlpA function without SlpX. How SlpX expression is upregulated is not clear yet. The promoter region of slpX in SB3 is identical to that of the parent. SlpX overexpression might be caused by transcriptional regulation responding to some (stress) condition when SlpB is expressed instead of SlpA, but clearly, the mechanism of overexpression of SlpX and the role of this minor SLP need further attention.

Previous studies have shown that some SLP or SLAP gene deletion strains of L. acidophilus, such as knockout strains of slpX and aggregation-promoting factor gene apf, altered bacterial stress tolerance [18, 34]. Our data showed that the stress tolerance of L. acidophilus L-92 expressing recombinant SlpAr1, SlpAr2, SlpAr4 or SlpB/X was changed compared with the parent strain. The strains expressing SlpAr1, SlpAr2 and SlpAr4 are more sensitive to sodium chloride, ox gall or ethanol, respectively. In addition, expression of recombinant SlpAr2 or SlpB/X affected SLAP protein patterns. The data suggest that SLPS play a crucial role in growth under certain stress conditions either directly or

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**Fig. 5.** In vitro binding of L. acidophilus strains to host proteins. An enzyme-linked immunosorbent assay (ELISA) was used to measure binding of the cells with DC-SIGN-Fc or uromodulin (Umod)-Fc chimera proteins. Heat-killed exponential-phase cells were coated on 96-well plates. DC-SIGN-Fc or Umod-Fc was added to the plates. Bound protein was detected with a horseradish peroxidase (HRP)-conjugated goat anti-human IgG-Fc fragment antibody and tetramethylbenzidine (TMB) by measuring absorbance. Data are the means of triplicate wells. The experiment was repeated three times with similar results. Error bars indicate the standard deviation. Statistical significance was calculated between the parent strain and each variant strain (Dunnett test). *p<0.05; **p<0.01; ***p<0.001.
by changing SLAP patterns on the surface of the cells. SLPs and SLAPs might function as barriers against detergents, alcohols or salts, all of which are known to have an influence on cell envelope integrity \cite{35}. Interestingly, stress sensitivity depends on the specific SLP expressed, namely SlpA or SlpB/X. Stress tolerance is also dependent on the position of the c-myc epitope in SlpA. Expression of SlpB/X or recombinant SlpA instead of wild-type SlpA could change the lattice monolayer structure of self-assembled SLPs, which could affect the robustness of SLP assembly, the strength of its binding to the cells, the barrier function, and/or the pattern and amount of SLAPs. One or a combination of these presumed changes might affect the bacterial stress tolerance comprehensively. The tertiary and quaternary structures of SLPs are not known. A better understanding of these structures and the molecular mechanisms of stress tolerance functions of SLPs and SLAPs should allow for the development of highly stress-tolerant strains.

Binding of the host proteins uromodulin and DC-SIGN by \textit{L. acidophilus} has been postulated to take place via SlpA \cite{19, 24}. Therefore, it was anticipated that the ability to bind these proteins of the strains expressing SlpA versions with a surface-exposed c-myc epitope, SAr1 and SAr2, might have changed. Here we showed that the binding of uromodulin and DC-SIGN is not only altered in strains SAr1 and SAr2, might have changed. Here we showed that the binding of uromodulin and DC-SIGN is not only altered in strains SAr1 and SAr2, but is also altered in strain SAr4, producing an SlpA variant with a non-surface-exposed c-myc epitope. This suggests that this recombinant version of SlpA might still affect the structure of the SLP monolayer and/or the pattern of SLAPs in such a way as to result in an altered binding of these host proteins. The binding of DC-SIGN and uromodulin to strain SAr1 was decreased to 79% and 50% of that of the parent strain, respectively, and this was quite similar to the decrease in binding to strain SAr4 (70% in each case). Whether these alterations in binding are a direct effect on possible contact sites between the human proteins and SlpA or due to changes in the general structure of the surface layer will need further clarification. The \textit{in vitro} binding of DC-SIGN to the SlpB- and SlpX-expressing strain SB3 did not change, but binding was significantly increased for uromodulin, suggesting that the binding of the 2 host proteins to surface layer proteins is not specific to SlpA. The 2 host proteins likely bind to SlpB and/or SlpX, the SLPs expressed instead of SlpA in strain SB3. Our data also suggest that more uromodulin binds to SlpB or SlpX than to SlpA and that the parts of surface layer proteins that make contact with the 2 host proteins are different. Strain SB3 might be efficiently incorporated into M cells via its binding ability to uromodulin, which could lead to effective modification of the host immune system. If proteins such as SlpB or SlpX allow for the producer cells to be efficiently incorporated into M cells, they might be applied for the future development of oral immune-modulating vehicles, such as recombinant bacteria, microcapsules or liposomes carrying these proteins on the surface. Whether SlpB and/or SlpX can bind to uromodulin strongly enough for the proteins to be efficiently delivered through M cells, which could lead to further efficient immune-modulation strategies, still needs to be elucidated.

Various reports exist on factors in \textit{L. acidophilus} modifying the host immune system. A lipoteichoic acid (LTA)-deficient \textit{L. acidophilus} mutant, obtained by deleting the phosphoglycerol transferase gene LAB0447, reduced IL-12 and enhanced IL-10 production in DCs, resulting in anti-inflammatory signalling.

\textbf{Fig. 6.} \textit{In vitro} cytokine production from dendritic cells (DCs) incubated with \textit{L. acidophilus} cells. Immature DCs were incubated with heat-killed \textit{L. acidophilus} cells expressing (recombinant) SlpA. Concentrations of the indicated cytokines in the supernatants of the DCs were determined using Bio-Plex system. Data are the means of values relative to the cytokine levels in DC culture with the \textit{Aupp} strain from two independent experiments performed in duplicate. Error bars indicate the standard deviation. Statistical significance between the data of the parent strain and each mutant was calculated using 2-way ANOVA and the post hoc paired t-test with Bonferroni correction. *p<0.05.
expression vector for mUmod-Fc.

Center for Integrative Medical Sciences) for kindly providing an

have been affected by changes in SLPs, as described above.

directly by the SLP changes or indirectly via SLAPs, which might

strains presented here might have been caused

cytokine production in DCs co-cultured with the recombinant

SLAPs might be different between the SlpB-expressing NCFM

cultured with the mutant. The amounts and patterns of SlpX and

increased pro-inflammatory cytokine production from DCs co-

of DC-SIGN to strain SB3 expressing SlpB, nor did we detect

from co-cultured DCs [19]. We did not detect decreased binding

pro-inflammatory cytokines such as IL-12p70, TNFα and IL-1β

change DC signaling. In a previous report, an SlpB expressing

results indicate that a recombinant form of SlpA, SlpAr2, might

showed that

an

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LTA is thought to be one of the main factors

and active cutaneous anaphylaxis as well as 2,4-dinitroflurobenzene and mine fetal

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