Abstract: *Streptococcus suis* is a Gram-positive bacterium, which causes sepsis and meningitis in pigs and humans. This review examines the role of known *S. suis* virulence factors in adhesion and *S. suis* carbohydrate-based adhesion mechanisms, as well as the inhibition of *S. suis* adhesion by anti-adhesion compounds in *in vitro* assays. Carbohydrate-binding specificities of *S. suis* have been identified, and these studies have shown that many strains recognize Galα1-4Gal-containing oligosaccharides present in host glycolipids. In the era of increasing antibiotic resistance, new means to treat infections are needed. Since microbial adhesion to carbohydrates is important to establish disease, compounds blocking adhesion could be an alternative to antibiotics. The use of oligosaccharides as drugs is generally hampered by their relatively low affinity (micromolar) to compete with multivalent binding to host receptors. However, screening of a library of chemically modified Galα1-4Gal derivatives has identified compounds that inhibit *S. suis* adhesion in nanomolar range. Also, design of multivalent Galα1-4Gal-containing dendrimers...
has resulted in a significant increase of the inhibitory potency of the disaccharide. The
*S. suis* adhesin binding to Galα1-4Gal-oligosaccharides, Streptococcal adhesin P (SadP),
was recently identified. It has a Galα1-4Gal-binding N-terminal domain and a C-terminal
LPNTG-motif for cell wall anchoring. The carbohydrate-binding domain has no homology
to *E. coli* P fimbrial adhesin, which suggests that these Gram-positive and Gram-negative
bacterial adhesins recognizing the same receptor have evolved by convergent evolution.
SadP adhesin may represent a promising target for the design of anti-adhesion ligands for
the prevention and treatment of *S. suis* infections.

**Keywords:** bacterial adhesion; galabiose; virulence; *Streptococcus suis*, carbohydrate;
adhesin; Gb3; galactose

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1. **Introduction**

*Streptococcus suis* is a Gram-positive bacterium, which is an emerging cause of serious infections,
such as meningitis, septicemia, endocarditis and pneumonia in pigs and zoonotic meningitis in
humans [1–4]. There are currently 35 known capsular serotypes in *S. suis*, of which the serotype 2
capsule (SS2) is the most virulent. Recently, multilocus sequencing has been used to more accurately
identify the relatedness of virulent strains. Based on this, *S. suis* sequence type 1 and 7 (ST1 and ST7)
have been found to be associated with the most severe infections [5].

Host- and tissue-specific adhesion of both the Gram-negative and Gram-positive bacteria is a
prerequisite for infection and invasive disease [6,7]. Bacteria interact with host mucosal cells in order
to colonize their ecological niche. Adhesion is important for avoiding the cleaning mechanisms of the
host mucociliary system. Bacteria have evolved multiple adhesins to specifically recognize host cell
surface carbohydrate and protein receptors [8].

*S. suis* has rapidly evolved drug resistance against antibiotics by horizontal transfer [9]. Therefore,
novel means to prevent and treat infections are needed. Anti-adhesion therapy is based on the
inhibition of bacterial attachment to a specific receptor structure [10,11]. Once the structures of the
receptors are known, more potent high-affinity receptor analogs can be designed. An advantage of the
prevention of infections by receptor analogs could be that bacteria do not develop resistance, in
contrast to the traditional bactericidal drugs. Since anti-adhesive compounds target adhesins, which are
required for a specific colonization of the host, the mutant phenotypes of bacteria deficient in adhesion
would be eliminated by the host.

Previous studies have shed new light on how *S. suis* interacts with host cells. In many cases,
virulence genes are involved directly or indirectly with adhesion. In this review, virulence mechanisms
with a role in adhesion are discussed. Particularly, *S. suis* adhesion mechanisms that are based on the
recognition of carbohydrate receptors, as well as the development of carbohydrate-based anti-adhesive
compounds are reviewed.
2. *Streptococcus suis* Adhesion

*Streptococcus suis* of the capsular polysaccharide type 2 is globally the most common isolate from both the porcine and human cases of meningitis [12]. It is typically isolated from the nasal cavity and tonsils of the pigs, which can carry this bacterium asymptotically for a long period of time. The exact mechanisms, which trigger the infectious disease and by which *S. suis* invades the blood circulation, are not known. It is thought that *S. suis* colonizing the upper respiratory tract may invade through the respiratory epithelium or via the intestine [13]. Once the bacteria have reached the blood circulation, they may penetrate into the brain by adhering to brain capillary endothelial cells and cross the blood brain barrier. The molecular mechanisms of *S. suis* interaction with different tissue compartments and cells are poorly known.

*S. suis* adhesion and invasion to the cultured epithelial cells commonly used in laboratories, such as A549, HeLa, MDCK and Hep-2 cells, has been studied. A polysaccharide capsule has been found to partially mask the adhesion [14]. The unencapsulated serotype 2 and non-typeable strains have been found to be more adhesive and invasive than encapsulated strains [15]. The invasion mechanisms of *S. suis* into host cells have been suggested to be closely related to other *Streptococci*. *S. suis* was described upon interaction with the host cell membrane to form large membrane invaginations or filopodia-like protrusions, followed by a process called close-contact-induced membrane-triggering [15].

Adhesion to the capillary endothelial cells precedes the penetration of *S. suis* into the brain from blood circulation and could be mediated by either direct invasion of bacteria into the cells by endocytic mechanisms or by disrupting the cell junctions. The adhesion of *S. suis* to the brain microvascular cells has been reported to be independent of the capsular polysaccharide expression. *S. suis* type 2 strains have been found to adhere to both human and porcine microvascular endothelial cells, but to invade only porcine cells with actin-dependent mechanisms [16]. Recently, *S. suis* and *Neisseria meningitidis* have been found to invade into the human choroid plexus papilloma cells and cross from the basolateral onto the apical side, thus modelling bacterial entry from the blood into the cerebrospinal side [17,18]. *S. suis* can also bind and activate plasminogen to proteolytically make the blood-brain barrier leaky to bacteria [19]. Furthermore, the *S. suis* polysaccharide capsule and cell wall can synergistically induce production of prostaglandin and metalloproteinase by macrophages and play a critical role in the permeabilization of the blood-brain barrier [20].

3. Virulence Genes with a Role in Adhesion

3.1. Regulators

During the colonization of the upper respiratory tract and the invasion of the blood circulation, *S. suis* needs to regulate the expression of the proteins involved in adhesion. Recent evidence shows that regulators required for *in vivo* virulence (Table 1) also modulate adhesion properties.

The regulatory genes required for virulence have been characterized by the comparison of the knockout mutants and wild-type bacteria for their survival both in mice and in pig infection models. Several two-component regulatory systems (TCS) regulate gene expression in *Streptococci*, such as those required for the natural competence of genetic transformation and quorum sensing [21,22]. TCS consist of a sensor kinase protein, which phosphorylates and activates a transcriptional regulator that
binds to promoters to induce or repress the expression of a large number of genes [23]. An orphan two-component transcriptional regulator can also control the transcription of virulence genes. Knockout strains harboring deletion in the *S. suis ciaRH* [24] and in the genes *revS* and *revSC21* encoding orphan regulators show decreased adhesion to Hep-2 cells [25,26]. SalK/SalR is another recently described TCS required for the virulence of *S. suis* in the porcine model, but its role in the regulation of adhesion is not well known [27,28]. The two-component system CovR/S regulates a number of *S. pyogenes* virulence genes. *S. suis* has an orphan orthologue of *covR*, also known as *csrR* (capsule synthesis regulator), that was found to negatively modulate virulence [29]. *S. pyogenes* Rgg, also known as RopB, is a transcriptional regulator known to regulate the expression of dozens of genes and has been shown to repress adhesion to epithelial cells and negatively regulate virulence in a murine intraperitoneal model [30–33]. In *S. suis*, the genes, *covR* and *rgg*, are required for virulence, and the corresponding knockout strains show increased binding to Hep-2 cells [29,34].

**Table 1.** *S. suis* virulence genes with known phenotype in adhesion or interaction with host cells. The chromosomal locations of the genes in the genome of *S. suis* type 2 P1/7, GZ1 or ST1 [35] are indicated.

| GENE ID       | GENE DESIGNATION | MUTANT PHENOTYPES                              |
|---------------|------------------|------------------------------------------------|
| SSU0944,      | *ciaRH* [24]     | Two-component regulator, decreased adhesion to Hep-2 cells |
| SSU0945       |                  |                                                 |
| SSU1873 not functional in P1/7, | *revS* [25]     | Orphan regulator, decreased adhesion to Hep-2 cells |
| SSGZ1_1897 in *S. suis* GZ1 |                  |                                                 |
| –             | *luxS* [36]      | Quorum sensing regulator, decreased adhesion |
| SSU0376       |                  |                                                 |
| SSU1789       | *rgg-like regulator* [34] | Negative transcriptional regulator, increased adhesion to Hep-2 cells |
| SSU1191       | *covR* [29]      | Orphan regulator, increased adhesion to Hep-2 cells |
| SSU1202       | *ccpA* [37]      | Carbon catabolite protein, decreased capsule thickness |

**Modulators of adhesion**

| GENE ID       | GENE DESIGNATION | MUTANT PHENOTYPES                              |
|---------------|------------------|------------------------------------------------|
| SSU0516,      | *cps2B, cps2E,   | Polysaccharide synthesis genes, increased adhesion of unencapsulated |
| SSU0519,      | *cps2F* [15,38], |                                                   |
| SSU0520,      | *cps2C* [39],    |                                                   |
| SSU0517,      | *neuB* [39]      |                                                   |
| SSU0535       |                  |                                                   |
| SSU0596,      | *dltA and pgdA* [40,41] | Cell wall modification, upregulated upon contact with endothelial cells |
| SSU1448       |                  |                                                   |

**Moonlighting or other cell wall proteins without signal sequence and known anchoring mechanism**

| GENE ID       | GENE DESIGNATION | MUTANT PHENOTYPES                              |
|---------------|------------------|------------------------------------------------|
| SSU0187,      | *dpp4* [42]     | Dipeptidyl peptidase IV, fibronectin binding |
| SSGZ1_0184    |                  |                                                 |
| SSU1320       | *eno* [43,44]    | Enolase, fibronectin and plasminogen binding, recombinant protein |
| SSU0153       | *GAPDH*, [45]    | Recombinant protein inhibits bacterial binding to porcine tracheal rings and Hep-2 cells |
| SSU1541       | *gnd* [46]       | 6-Phosphogluconate-dehydrogenase, recombinant protein inhibits bacterial binding to Hep-2 and HeLa cells |
| SSU0157       | *glnA* [47]      | Glutamate synthetase, decreased adherence to the Hep-2 cells |
| SSU1127       | *atl, autolysin* [48] | Biofilm and Hep-2 cell adhesion |
| SSU1311       | *fbps* [49]      | Fibronectin binding |

Table 1. *S. suis* virulence genes with known phenotype in adhesion or interaction with host cells. The chromosomal locations of the genes in the genome of *S. suis* type 2 P1/7, GZ1 or ST1 [35] are indicated.
Table 1. Cont.

| GENE ID   | GENE DESIGNATION | MUTANT PHENOTYPES                                                   |
|-----------|------------------|-------------------------------------------------------------------|
| SSU0925   | srtA [50]        | Anchoring of cell wall proteins                                   |
| SSU0879   | IgA1 protease [51,52] | Degradation of IgA protecting mucosal surfaces                    |
| SSU1474 (pseudo), SSUST1_1540 in S. suis ST1 | sof [53] | Lipoprotein degradation                                           |
| SSU0757   | sspA, [54,55]    | Subtilisin-like protease, induces secretion of cytokines and chemokines |
| SSU1143   | ssa [56]         | Fibronectin/fibrinogen binding, reduced adhesion and invasion to Hep-2 cells |

An autoinducer AI-2/LuxS quorum sensing system regulates virulence and biofilm formation in Streptococci [57,58]. In S. suis, it positively regulates biofilm formation and adhesion to Hep-2 cells [36]. In addition, a deletion mutant in the gene, stp, which encodes a serine/threonine phosphatase, has been shown to adhere less to Hep-2 cells compared to the wild-type strain [59]. A global regulatory protein, called catabolite control protein, CcpA, regulates the utilization of carbohydrates by S. suis. It also regulates polysaccharide capsule thickness and hemolytic activity [37]. The regulation of S. suis capsule thickness might affect the adhesion properties. Conclusively, in S. suis, a complex network of regulation of virulence also modulates adhesion.

3.2. S. suis Surface Glycoconjugates

The S. suis capsular polysaccharide can block the adhesins at the cell surface. Unencapsulated strains agglutinate erythrocytes and adhere more strongly to HEp-2 cells than encapsulated strains [15,60]. The serotype 2 polysaccharide structure consists of the repeating unit [(Neu5Acα2-6Galβ1-4GlcNAcβ1-3)Galβ1-4(Galα1-3)Rhaβ1-4Glcβ1-]n [61]. Presence of sialic acid in S. suis capsular polysaccharide creates a negative charge on the bacterial surface. Therefore, it is likely that a host cell surface negative charge and the sialic acid containing bacterial polysaccharide form a biophysical repulsion strong enough to reduce adhesion of encapsulated S. suis to host cells. Recently, mutant strains with the deletion of genes required for serotype 2 capsular polysaccharide synthesis were found to be avirulent in animal models, but more adherent and invasive to Hep-2 cells [39].

Streptococcus agalactiae polysaccharide contains sialic acid and is important for the evasion of the complement and phagocytosis, but in the case of S. suis, it has been suggested that, due to the lower amount of sialic acid in S. suis type 2 capsule, it does not have a major role in the evasion of the complement [61]. Instead, the polysaccharide capsule of type 2 was previously described to be involved in the interaction of S. suis with macrophages. Specifically, removal of S. suis sialic acid reduced its binding to J774 macrophages [62]. The ligands recognizing sialic acid are not known. The S. suis type 2 capsular polysaccharide was recently found to prevent phagocytosis of S. suis by destabilizing lipid microdomains of phagocytes. The capsular polysaccharide prevented lactosylceramide accumulation into the phagocytic cup and the activation of lactosylceramide-lipid raft signaling pathways, thus inhibiting the activation of the phosphoinositide 3-kinase/Akt and p38 MAPK pathways required for the activation of phagocytosis [63]. The ability of S. suis to adhere to immune
cells and to prevent phagocytosis could be a mechanism for the bacteria to cross the blood-brain-barrier, as suggested by the “modified Trojan horse” theory [63].

The role of the \textit{S. suis} polysaccharide capsule in the invasion of bacteria into the brain has been studied in an \textit{in vitro} model of neutrophil transmigration. The neutrophils infected with \textit{S. suis} transmigrated through the choroid plexus epithelial cell monolayer in an inverted Transwell system mimicking the blood-brain barrier [64]. Encapsulated \textit{S. suis} bacteria induced the granulocytes to transmigrate through the porcine choroid plexus epithelial cells from their basolateral sides. This correlates with the theory that the influx of leukocytes into the brain is important for the development of meningitis.

Mutant strains harboring deletions in the alanine-alanine ligase, \textit{dltA}, and peptidoglycan \textit{N}-deacetylase \textit{pgdA} genes, required for modification of lipoteichoic acid and peptidoglycan structure, are more susceptible to the action of antimicrobial peptides and neutrophil killing mechanisms [40,41]. In addition, these genes are required for \textit{in vivo} survival in animal models. Interestingly, the transcription of alanine-alanine ligase and peptidoglycan deacetylase were upregulated upon contact with endothelial cells [65]. It was suggested that the regulation of these genes can prime bacteria to be more resistant for neutrophil killing after they have crossed the blood-brain barrier.

\subsection*{3.3. Cell Wall Proteins}

\textit{S. suis} have evolved mechanisms to adhere to the host cells utilizing moonlighting proteins, \textit{i.e.}, cell wall-attached proteins that have enzymatic activities in addition to adhesion specificities [67]. Several \textit{S. suis} enzymes have been reported to be adhesins. Except for interactions with host fibronectin, the host receptors have remained unknown. Fibronectin has been found to be recognized by the \textit{S. suis} cell wall-bound proteins, dipeptidyl peptidase IV, enolase and Fbps [42–44,49]. Enolase recognizes also plasminogen. Plasminogen binding, in addition to fibronectin, could cooperatively help to degrade extracellular matrix and fibrin and aid in the invasion of bacteria into tissues. Plasminogen binding seems in many \textit{Streptococci} to be important in invasion [68]. The adhesion properties of moonlighting proteins and their ability to activate host proteases might represent an example of convergent evolution of bacteria to be able to use host proteases in subverting host defense mechanisms. Furthermore, \textit{S. suis} glyceraldehyde-3-phosphate dehydrogenase, 6-phosphogluconate dehydrogenase, glutamine synthetase and autolysin have been identified as moonlighting adhesins [45–48,69], but their adhesion specificities are not known.

LPXTG-motif containing proteins are covalently anchored to the cell wall peptidoglycan via their \textit{C}-termini. They require a sortase enzyme that covalently attaches them to the peptidoglycan. \textit{S. suis} housekeeping sortase A (SrtA) seems important in adhesion, since a knockout strain was shown to be
deficient in the adhesion to endothelial cells [50]. Recently, \textit{srtA} was shown to be required for high virulence and adhesion to Hep-2 cells and human umbilical endothelial cells [70]. Some of the LPXTG-anchored proteins, such as IgA protease, serum opacity factor Sof, subtilisin-like protein SspA and fibronectin-binding protein Ssa are known to be required for virulence [51–56]. Several other LPXTG-motif proteins have been described, but relatively little is known of their role in adhesion and virulence (for extensive reviews, see [13,71]).

An important group of proteins containing the LPXTG-anchor motif are pilins, which form the fimbrial structures, called pili, in \textit{Streptococci} [66]. Typical for the fimbriae of Gram-positive bacteria are that the pilins are covalently linked via motifs closely resembling the LPXTG sequences. The linkages are catalyzed by pilus islet-specific sortases. In \textit{S. suis}, the pilus islands srtBCD, srtE, srtF and srtG have been identified [72,73]. The SrtF cluster contains four genes encoding a putative signal peptidase (\textit{siF}), putative ancillary (\textit{sfp2}, pseudogene) and major (\textit{sfp1}) pilus subunits and class C sortase, \textit{srtF} [74]. Surprisingly \textit{sfp2}, which is homologous to other Streptococcal fimbriae-associated adhesins, is a pseudogene, and the experiments carried with WT and \textit{sfb1} mutants indicate that \textit{S. suis} fimbriae are indispensable for attachment to the pig brain capillary endothelial cells and for virulence in mouse sepsis models [74].

Interestingly, studies characterizing gene expression upon bacterial contact with endothelial cells have revealed that the mRNA levels of signal peptidase and sortase E involved in pilus synthesis were increased [65]. This suggests that, with the so far unknown mechanisms, \textit{S. suis} can sense the contact with host cells and induce the expression of genes important for interaction with host cells.

4. \textit{S. suis} Carbohydrate-Specific Adhesion to Host Cells

A limited number of \textit{S. suis} carbohydrate-specific adhesion mechanisms have been described. Some \textit{S. suis} strains recognize sialic acid-containing oligosaccharides. Bacteria bind to the terminal sialic acid in polylactosamine chains with the fine specificity of NeuN\textit{Ac}a2-3Gal\beta1-4Glc\textit{NAc}β1-3Gal [75]. Recently, two \textit{S. suis} LPXTG-anchored proteins have been shown to recognize host cell carbohydrates. One of them, protein HP0197, binds host cell surface glycosaminoglycans. The adhesin knockout strain had decreased binding to Hep-2 cells [76]. We have previously identified a Galα1-Gal-binding activity and identified, recently, the corresponding adhesin [77,78]. The binding specificity to Galα1-4Gal (galabiose) and exploitation of receptor analogs in the development of anti-adhesion compounds are discussed below.

4.1. Galabiose (Galα1-4Gal)-Specific Adhesion of \textit{S. suis}

The \textit{S. suis} galabiose-specific adhesion activity was originally characterized with hemagglutination assay based on multivalent recognition of erythrocyte surface carbohydrates by bacterial adhesins [77]. Galα1-4Gal-oligosaccharides are present in globo-series glycolipids, which are abundant in erythrocytes, endothelial and uroepithelial cells. Glycolipids containing the Galα1-4Gal-sequence form the human P blood group system. The \textit{S. suis} galabiose binding strains can be grouped into two groups based on their monosaccharide inhibition pattern in hemagglutination assays [77]. Type \(P_N\) is inhibited by both galactose and \(N\)-acetylgalactosamine, whereas type \(P_O\) is only inhibited by galactose. The \textit{S. suis} hemagglutination is inhibited with oligosaccharides containing the Galα1-4Gal structure at micromolar
concentrations, but only at millimolar concentrations by disaccharides containing Gal\(\alpha_1\)3Gal or Gal\(\alpha_1\)6Glc linkages [77]. Analysis of the detailed adhesion specificity toward galabiose-containing glycans revealed that it has the highest specificity to globotriaosylceramide GbO3, Gal\(\alpha_1\)4Gal\(\beta_1\)-4Glc\(\beta_1\)1-1'Cer (the \(P^k\) antigen, Table 2).

**Table 2.** Gal\(\alpha_1\)-4Gal-binding proteins \(^a\).

| Structure/Antigen \(^a\) | Ligands |
|-------------------------|---------|
| **GbO3 \(^b\)** | Gal\(\alpha_1\)-4Gal\(\beta_1\)-4Glc\(\beta_1\)-1'Cer/\(P^k\) |
| GbO4 | GalNAc\(\beta_1\)3Gal\(\alpha_1\)-4Glc\(\beta_1\)-1'Cer/P |
| GbO5 | GalNAc\(\alpha_1\)-3Gal\(\alpha_1\)-4Gal\(\beta_1\)-4Glc\(\beta_1\)-1'Cer/Forssman |

**Adhesins**
- *E. coli* PapGI [79], *S. suis* SadP, *P. aeruginosa* Lectin I [80]

**Toxins**
- *S. dysenteriae* Shiga toxin [81], *E. coli* verotoxin 1, 2, 2c [82,83]
- *E. coli* verotoxin 2e [84]

\(^a\) For simplicity, blood group antigen P\(^1\), **Gal\(\alpha_1\)-4Gal\(\beta_1\)-4Glc\(\beta_1\)-1'Cer/\(P^k\)**, is not included in the table; \(^b\) GbO3, GbO4 and GbO5 are abbreviations for globotria-, globotetra- and globopentaosylceramides.

Gal\(\alpha_1\)-4Gal is also recognized by *E. coli* PapG adhesins, *Pseudomonas aeruginosa* lectin I, Shiga toxin and verotoxins (Table 2). The globo-series glycolipid composition varies between animal species, and different galabiose containing glycolipids are host-specifically recognized by uropathogenic *E. coli* P fimbrial adhesins [79] and *E. coli* verotoxins (Table 2).

4.2. The Galabiose-Binding Adhesin SadP

The *S. suis* galabiose-specific adhesin has been isolated [78]. The adhesin was extracted from the *S. suis* cell wall with lysozyme and was captured with the pigeon ovomucoid affinity matrix. Pigeon ovomucoid is a strong glycoprotein inhibitor of *S. suis* hemagglutination, as it contains terminal Gal\(\alpha_1\)-4Gal-residues in its oligosaccharides [85,86]. A gene encoding the adhesin was identified as SSU_0253 from the *S. suis* serotype 2 strain P1/7 genomic sequence. Sequence analysis of the gene shows that it has an N-terminal signal peptide for secretion, seven C-terminal tandem repeats and an LPXTG-motif. An *S. suis* sadP knockout strain lost galabiose binding activity. A recombinant N-terminal domain of the adhesin recognizes Gal\(\alpha_1\)-4Gal-oligosaccharides. The adhesin was designated as the Streptococcal adhesin P (SadP).

The SadP galabiose-binding domain has no significant homology to other known galabiose-binding proteins, including *E. coli* P fimbrial adhesins and Shiga toxins and verotoxins (Table 2). An identical *S. suis* protein, Fhb (HP0272), has been reported to bind to human complement factor H, which is a glycoprotein that regulates the complement system [87]. The factor H binding domain is located in the C-terminal region, which has homology to *S. agalactiae* IgA-binding protein [78]. The factor H binding protein in *S. suis* strain 05ZYH33 has an N-terminal domain that is identical to SadP. The *fhb*-05ZYH33 knockout strain was highly attenuated in a piglet model [87].

In the galabiose structure, the essential key hydroxyls for *S. suis* binding are HO-4', HO-6', HO-2 and HO-3 (Figure 1), and in addition, P\(_0\) strains have weak interactions with HO-3' and HO-6. The
combining site of type $P_O$ is more narrow on the terminal position of galactose, whereas the $P_N$ strains allow substitutions in the HO-3' position. The essential hydroxyls of galabiose required for the binding of *E. coli* PapG adhesins are different. *E. coli* PapG recognizes HO-2', HO-3', HO-4', HO-6' and HO-6 hydroxyls in the disaccharide (Figure 1). The hydrogen bonding pattern suggests that these two bacterial adhesins recognize the disaccharide from different sides [77]. Conclusively, the functional assays with oligosaccharide derivatives and the adhesin sequence comparisons indicate that the galabiose binding *E. coli* PapG and *S. suis* SadP have different modes of mechanisms to interact with Gal$_\alpha$-4Gal and represent an example of convergent evolution of bacterial adhesins toward binding to the same Gal$_\alpha$-4Gal-receptor.

Figure 1. (a) Hydrogen bonding patterns of *S. suis* P$_N$, *S. suis* P$_O$ [77] and *E. coli* PapG adhesins [88,89]. Black dashed lines indicate hydrogen bonds and grey dashed lines indicate possible weak hydrogen bonds. Hydrogen bond directionalities have been mapped for the *E. coli* PapG class I adhesin [90], which was later confirmed by structural analysis of a globotetraose:papG class II adhesin complex [91]. (HBA = hydrogen bond acceptor, HBD = hydrogen bond donor). Derivatization with urea and ether groups at O-3 of $\alpha$-D-Gal [92] enhances affinity for *S. suis* P$_N$ and *E. coli* PapG, respectively (indicated with yellow ovals). Ether groups at O-2 of $\alpha$-D-Gal enhance binding to *S. suis* P$_N$ (indicated with green oval). (b) Space-filling models of the globotriose trisaccharide with hydroxyl oxygens interacting with the three adhesins shown in light pink, which illustrates the different epitopes recognized. (Globotriose conformation taken from the globotetraose:papG class II adhesin complex; pdb id 1J8R.)

5. Towards the Development of Therapy Based on Prevention of Adhesion

Carbohydrates have usually a low affinity to adhesins, partly because the $O$-glycosidic linkages between monosaccharides in oligosaccharides are very flexible. In addition, monovalent oligosaccharide inhibitors are inefficient for competing with the multivalent receptor interactions. Strategies to overcome the relatively low affinity of the lectin-carbohydrate interactions include modification of the hydroxyl groups of the oligosaccharide receptors and construction of multivalent carbohydrate dendrimers to increase the affinity [93,94].
5.1. Combinatorial Libraries of Receptor Carbohydrates

Previous data showed that \textit{S. suis} galabiose binding was not dependent on the HO-3' and HO-2' hydroxyls of the terminal α-galactose (Figure 1). The adhesin combining site of type P\textsubscript{N} allowed larger substituents in the HO-3' position of the terminal galactose, as compared to type P\textsubscript{O} adhesin. When large libraries of galabiose derivatives were tested for the inhibitory activity with \textit{E. coli} PapG and \textit{S. suis} SadP, galabiosides carrying aromatic structures at C1 were found to be efficient inhibitors of the PapG adhesin \cite{92}. The major difference between PapG and SadP was that methoxymethylation at O2' and phenylurea derivatization at C3' of the terminal galactose in the galabiose disaccharide provided inhibitors of the \textit{S. suis} type P\textsubscript{N} adhesin with nanomolar affinities.

5.2. Dendrimers as Polyvalent Carbohydrate Inhibitors

The mode of interaction between bacterial adhesins of \textit{S. suis} and \textit{E. coli} are different compared to Shiga-like toxins with GbO3 glycolipids. The adhesin can be inhibited with soluble oligosaccharides, whereas Shiga-like toxins are not inhibited and require that the receptor oligosaccharide is presented polyvalently. Utilizing the crystal structure of the AB5 Shiga-like toxin, a “STARFISH” oligomeric water-soluble compound was designed consisting of five trisaccharides at the tip of spacers. It was found to occupy all five receptor binding sites and had subnanomolar inhibitory activity \cite{95}. The inhibition of \textit{S. suis} type P\textsubscript{N} and P\textsubscript{O} hemagglutination activities with dendrimeric galabiosides have been studied \cite{96}. The inhibitions were tested under static conditions using hemagglutination in microtiter wells. The relative potencies of the dendrimeric compounds increased to 300–400-fold when galabiose was presented as a divalent compound.

Bacteria adhere often under share stress and in dynamic flow conditions; therefore, a surface plasmon resonance assay has been used to analyze bacterial binding to galabiose-BSA neoglycoprotein under flow conditions \cite{97}. The inhibitory activity of divalent, tetravalent and octavalent galabiose dendrimers were tested (Table 3). The highest relative inhibitory potency of 250 per sugar was obtained with the tetravalent compound as compared to the monovalent disaccharide. The potency per sugar of the tetravalent compound was twice that of the octavalent compound. For \textit{E. coli} expressing an adhesin belonging to the PapGI class, which recognizes terminal Galα1-4Gal, the inhibitory potency per sugar was highest for the octavalent compound, which suggests that the octavalent compound has more possibilities to interact with the tip-located fimbrial adhesins.

\textit{S. suis} adhesion has been successfully inhibited with galabiose-containing oligosaccharide derivatives, and SadP adhesin is thus a promising target for anti-adhesion therapy. Derivatives of galabiose, as well as polyvalent galabiose dendrimers can inhibit adhesion of \textit{S. suis} in the nanomolar range. These results suggest that anti-adhesion therapy could be a potential alternative to antibiotics in the treatment of \textit{S. suis} infections.
Table 3. Inhibitory potencies of galabiose dendrimers against *S. suis* and *E. coli* adhesion (modified from [98]).

| Bacterial strain and inhibitor | Valency of dendrimer | Relative potency | Potency per sugar |
|-------------------------------|----------------------|------------------|-------------------|
| *S. suis* 628 \(^a\)          |                      |                  |                   |
| Monovalent                    | 1                    | 1                | 1                 |
| Divalent, short spacer        | 2                    | 13               | 7                 |
| Divalent, long spacer         | 2                    | 12               | 8                 |
| Tetravalent                   | 4                    | 250              | 63                |
| Octavalent                    | 8                    | 310              | 39                |
| Octavalent PAMAM              | 8                    | 260              | 32                |
| *S. suis* D282 \(^b\)         |                      |                  |                   |
| Monovalent                    | 1                    | 1                | 1                 |
| Divalent                      | 2                    | 50               | 25                |
| Tetravalent                   | 4                    | 170              | 42                |
| Tetravalent galatriose \(^c\) | 4                    | 8                | 2                 |
| Octavalent                    | 8                    | 100              | 13                |
| *E. coli* PapGJ96 \(^d\)      |                      |                  |                   |
| Monovalent                    | 1                    | 1                | 1                 |
| Divalent, short spacer arms   | 2                    | 3                | 1                 |
| Divalent, long spacer arms    | 2                    | 3                | 1                 |
| Tetravalent                   | 4                    | 8                | 2                 |
| Octavalent                    | 8                    | 43               | 5                 |
| Octavalent PAMAM              | 8                    | 6                | 1                 |

\(^a\) The IC\(_{50}\) values were determined in a surface plasmon resonance (SPR) assay [97]; relative potency = IC\(_{50}\) or the MIC of the monovalent inhibitor divided by that of the inhibitor in question; relative potency per sugar = relative potency/valency; \(^b\) MIC values were determined in a hemagglutination assay [99]; \(^c\) other inhibitors were galabiose derivatives.

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Conflicts of Interest

The author declares no conflict of interest.

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