1	Brachyspira hyodysenteriae binding to porcine colonic mucins differs between individuals
2	and is increased to mucins from infected pigs with de novo MUC5AC synthesis
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25 Abstract

26 Brachyspira hyodysenteriae colonizes the pig colon resulting in mucohemorrhagic diarrhea and growth retardation. Fecal mucus is a characteristic feature of swine dysentery; therefore we 27 investigated how the mucin environment changes in the colon during infection with B. 28 29 hyodysenteriae, and how these changes affect this bacterium's interaction with mucins. We 30 isolated and characterized mucins, the main component of mucus, from the colon of experimentally inoculated and control pigs, and investigated *B. hyodysenteriae* binding to these 31 mucins. Fluorescence microscopy revealed a massive mucus induction and disorganized mucus 32 33 structure in the colon of pigs with swine dysentery. qPCR and antibody detection demonstrated that the mucus composition of pigs with swine dysentery was characterized by de novo 34 expression of MUC5AC and increased expression of MUC2 in the colon. Mucins from colon of 35 36 inoculated and control pigs were isolated by two-steps of isopycnic density-gradient centrifugation. The mucin density was similar between control and inoculated pigs, whereas the 37 mucin quantity was five-fold higher during infection. B. hyodysenteriae bound to mucins in a 38 manner that differed between pigs and there was increased binding to soluble mucins isolated 39 from pigs with swine dysentery. The B. hyodysenteriae binding ability, in relation to the total 40 41 mucin content of mucus from sick vs. healthy pigs, increased seven-fold during infection. Together, the results indicate that B. hyodysenteriae binds to carbohydrate structures on the 42 mucins, as these differ between individuals. Furthermore, B. hyodysenteriae infection induces 43 44 changes to the mucus niche, which substantially increases the amount of *B. hyodysenteriae* binding sites in the mucus. 45

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49 Introduction

50 The gastrointestinal tract is lubricated by a continuously secreted mucus layer which can also act as a barrier against pathogens (1). The main components of the mucus layer are heavily 51 glycosylated gel-forming mucins. Mucin glycans can prevent enzymatic degradation of the 52 mucin protein core, and can also bind water, conferring viscoelasticity (2). Underneath the 53 54 mucus layer, transmembrane mucins on the mucosal epithelial cells provide barrier and reporting functions (3, 4). Mucins differ in their glycosylation and tissue distribution (5). Murine colonic 55 mucus has been shown to be rich in the MUC2 mucin, which is secreted by goblet cells and is 56 57 organized in a two-layered mucus system (6). The inner mucus layer is firmly attached to the epithelium, and gives rise to the loosely adherent outer layer (7). 58

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Mucins are a dynamic component of the mucosal barrier, and have been shown to undergo changes in response to intestinal infection and inflammation in mice (8, 9). Mucin glycan structures can bind bacteria, *e.g. Escherichia coli* and *Helicobacter pylori*, limiting colonization and access to the epithelial surface (3, 10-13). Mucin glycosylation can change during bacterial infection, and varies between individuals (14). To date it is unknown whether the large variability in mucin expression and mucin glycosylation arose by chance during evolution, or if the mucin species confer distinct response properties during infection.

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Brachyspira hyodysenteriae (B. hyodysenteriae) is a recognized swine pathogen, commonly associated with swine dysentery (SD). This anaerobic spirochete colonizes the large intestine of pigs resulting in mucohemorrhagic diarrhea. Ingestion of feces from inoculated pigs, as well as from asymptomatic carriers, is among the main sources of infection (15). SD is responsible for economic losses in the swine industry, posing a threat in countries where antimicrobials are banned for growth promotion, and challenging those where resistant strains have emerged (16-19). The presence of mucus in the feces is a characteristic feature of SD. Recently, colonic specimens from pigs with SD were shown to have increased immunohistochemical staining with an antibody against the human MUC5AC mucin and decreased staining with an antibody against the MUC4 mucin (20).

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B. hyodysenteriae pathogenesis is still surrounded by many uncertainties. The mechanisms 79 underlying the bacterial interactions with the colonic mucosal surface, or how the mucin 80 81 response exerted during infection is regulated, remain to be elucidated. Therefore, the overall aims of the present study were to investigate how the mucin environment changes in the swine 82 83 colon during infection with *B. hyodysenteriae*; if this bacterium binds to mucins, and if so, how 84 these changes affect binding. We identified that B. hyodysenteriae infection causes changes in mucus organization, mucin quantity, identity and expression profile, as well as in the mucin 85 86 binding ability of this bacterium.

87

88 Materials and Methods

89 Ethics statement

90 The animal experiments were approved by the ethical committee of the Faculty of Veterinary
91 Medicine, Ghent University (EC2012/01 and EC2013/147) and complied with all ethical and
92 husbandry regulations.

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94 Experimental inoculation and sample collection

Samples from a total of 15 pigs (Danish Large White × Piétrain) from two independent
inoculation experiments, 21 months apart, were included in the study (Table 1). The first

97 experiment included 6 control pigs and 6 inoculated pigs, and the second 8 control pigs and 8 inoculated pigs. The pigs from both inoculation experiments were 6-week old, came from two 98 99 different commercial farrowing to finishing farms in the Flanders region with no previous history of swine dysentery, belonged to different litters, and were fed the same commercial starter feed 100 (crude protein 17%, crude fat 6.09%, crude fiber 3.87%, crude ash 5.09%, phosphorus 0.49%, 101 methionine 0.43%, lysine 1.25%, calcium 0.61%, and sodium 0.23%; Lambers-Seghers, 102 Belgium). At their arrival, the pigs were confirmed negative for *B. hyodysenteriae* in rectal fecal 103 samples by culture and qPCR. The pigs were acclimatized for two weeks in order to recover 104 105 from transport stress and adapt to diet and housing changes. The pigs were fed twice per day and had ad libitum access to water. A total of 14 pigs were experimentally inoculated with B. 106 hvodysenteriae strain 8dII, isolated from a Belgian swine farm with a history of recent dysentery 107 problems. An inoculum of 10⁸ CFU/ml in brain heart infusion broth (BHI) (50 ml/pig) was 108 administered orally during three consecutive days, while 14 control pigs received 50 ml of sterile 109 BHI. From the 14 control pigs, 6 samples were randomly selected for use in this study (pigs A-110 F). From the first infection trial two out of six pigs developed SD, and from the second infection 111 trial three out of eight pigs developed SD (pigs 1-5). Samples from the four inoculated pigs that 112 113 did not develop SD in the first inoculation experiment were included in the study (pigs 6-9; Table 1). 114

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Infection was confirmed based on clinical signs of mucohemorrhagic diarrhea, and *B. hyodysenteriae* excretion in feces detected by qPCR in fecal samples obtained twice a week. The pigs were sacrificed at day 40 post-inoculation by anesthesia with a combination of xylazine at 4.4 mg/kg (Xyl-M 2%[®], VMD, Arendonk, Belgium) and zolazepam/tiletamine at 2.2 mg/kg (Zoletil[®] 100, Virbac, Carros, France), and final euthanasia by intracardial injection of a

formulation comprising embutramide, mebezonium iodide and tetracaine (T61[®], Intervet,
Brussels, Belgium) at 0.3 ml/kg.

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Midsection samples of the spiral colon with a size of 7×8 cm were obtained from the inoculated 124 and control pigs for mucin isolation. Fecal material was removed, and the tissues were rinsed 125 126 with phosphate buffered saline (PBS) and a protease inhibitor cocktail (Roche Diagnostics, Mannheim, Germany) before snap freezing and storage at -80°C. Smaller specimens were 127 carefully collected without disturbing the mucus layer (no washing) and immersed in 10-volumes 128 129 of fresh Carnov's methanol fixative (60% dry methanol, 30% chloroform, 10% glacial acetic acid), and embedded in wax for histology/immunohistochemistry. There were also samples 130 collected in RNAlater® (Life Technologies, Carlsbad, CA, USA) and kept at 4°C overnight, then 131 132 stored at -80°C for RNA extraction.

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134 Detection of B. hyodysenteriae in feces by qPCR

DNA from pig feces was obtained by using the QIAamp DNA stool mini kit (Qiagen, CA, USA)
starting from 1 gram of feces. For qPCR, *Brachyspira* spp. specific primers were used in
combination with a *B. hyodysenteriae* specific probe as previously described (21).

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139 MUC2 and MUC5AC immunofluorescence

Tissue sections were deparaffinized and antigen retrieval was performed in 10 mM sodium
citrate, pH 6.0 at 99°C for 30 min. Slides were cooled to room temperature and washed in PBS.
Non-specific background was blocked with serum-free protein block (DAKO, Carpinteria, CA,
USA) for 20 min. Primary antibodies anti-MUC2C3 (kindly provided by G. Hansson, University
of Gothenburg, Sweden), anti-MUC5AC (45M1, Sigma-Aldrich, St. Louis, MO, USA) and anti-

145 MUC5CR (kindly provided by G. Hansson, University of Gothenburg, Sweden) were diluted 146 1/1000 and incubated at 4°C overnight. Sections were washed with PBS and incubated with secondary antibodies conjugated with Alexa Fluor 488 (Life Technologies, Eugene, OR, USA) 147 for MUC2, and Alexa Fluor 594 for MUC5AC, diluted 1/500 for 1 h. After washing in PBS, 148 149 specimens were mounted with ProLong® antifade containing DAPI (Life Technologies, Eugene, 150 OR, USA). Pig and human gastric specimens were used as positive controls for MUC5AC staining. Similarities in the stomach and colon binding patterns indicated the staining of pig 151 sections was specific, even though the antibodies were raised against human mucins. 152

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154 *qPCR for mucin expression*

Pig colon tissue samples were immediately submerged in a 10-fold volume of RNAlater® (Life 155 156 Technologies, Carlsbad, CA, USA) at 4°C overnight, and frozen at -80°C until RNA extraction. Isolation of RNA was performed using Trizol (Life Technologies, Carlsbad, CA, USA) 157 according to the manufacturer's instructions. RNA yield and purity was assessed through UV 158 spectroscopy (NanoDrop, Thermo Scientific, MA, USA). Total RNA (5 µg) was DNase treated 159 at 37°C for 45 min, followed by the addition of 5 mM ethylenediaminetetraacetic acid (EDTA) 160 161 and heat inactivation of DNase at 75°C for 10 min prior to cDNA synthesis. MgCl₂ was added to 162 a 5 mM final concentration, and this RNA was used for cDNA synthesis with random hexamers and Superscript III (Life Technologies, Carlsbad, CA, USA) at 50°C for 2 h. The cDNA was 163 164 used in a real-time PCR reaction using SYBR green (Power SYBR® green mix, Life Technologies, Carlsbad, CA, USA) and primers listed in Table 2. Primers for pig MUC1, MUC2, 165 and MUC5AC mucin genes were designed using the Primer3 program (available at 166 167 http://frodo.wi.mit.edu/primer3/). qPCR data were normalized using the expression levels of ACTB and RPL4 reference genes (22). Samples were amplified in triplicate, and a negative 168

169 control without reverse transcriptase was included to verify the absence of contaminating
170 genomic DNA. Data acquisition and analysis was performed using the CFX manager 3.1
171 software (Bio-Rad Laboratories Inc., Hercules, CA, USA).

- 172
- 173 Mucin isolation and purification

Mucin isolation of colon tissue samples was performed by isopycnic density gradient 174 centrifugation as previously described (23), obtaining guanidinium hydrochloride (GuHCl) 175 soluble and insoluble mucins. Briefly, frozen tissues were drenched with 10 mM sodium 176 177 phosphate buffer, pH 6.5, containing 0.1 mM phenylmethylsulfonyl fluoride (AppliChem, 178 Darmstadt, Germany). Once thawed, the mucosal surfaces were scraped with a microscope slide, dispersed with a Dounce homogenizer, and stirred slowly overnight at 4°C in ice-cold extraction 179 180 buffer consisting of 6 M GuHCl (AppliChem, Darmstadt, Germany), 5 mM EDTA (Sigma-Aldrich, St. Louis, MO, USA), 5 mM N-ethylmaleimide (Alfa Aesar, Karlsruhe, Germany) and 181 182 10 mM sodium di-hydrogen phosphate at pH 6.5. GuHCl soluble mucins were obtained after centrifugation at $23000 \times g$ for 50 min at 4°C and the remaining material was re-extracted twice 183 by stirring overnight at 4°C in extraction buffer. The remaining pellets contained the "insoluble" 184 mucins, which were solubilized with 10 mM dithiothreitol (DTT) in reduction buffer (6 M 185 GuHCl, 5 mM EDTA, 0.1 M Tris/HCl, pH 8) for 5 h at 37°C. Finally, residues were alkylated 186 overnight with 25 mM iodoacetamide (IAA, Alfa Aesar, Karlsruhe, Germany). 187

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Both the GuHCl soluble and insoluble material was dialyzed in ten volumes of extraction buffer at 4°C, changing the dialysis solution three times in 24 h. An isopycnic density-gradient centrifugation in cesium chloride (CsCl)/4 M GuHCl with a starting density of 1.39 g/ml was performed at 40000 rpm for 90 h. The mucin containing fractions were pooled and further purified from DNA by a second gradient in CsCl/0.5 M GuHCl. Approximately 25 mucin
fractions were recovered per sample using a fraction collector equipped with a drop counter.
Fractions were stored at 4°C until further analysis.

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197 Analysis of mucin fractions

First and second CsCl gradient mucin fractions were analyzed as follows. The mucin density was 198 determined by weighing a known volume using a Carlsberg pipette as a pycnometer, results were 199 expressed as g/ml. DNA contamination of mucins were determined using a spectrophotometer. A 200 201 microtiter-based assay detecting carbohydrates as periodate-oxidizable structures (24) was performed in order to determine the glycan content in the GuHCl soluble and insoluble mucin 202 samples. Briefly, Nunc® 96-well plates (Thermo Scientific, Waltham, MA, USA) were coated 203 204 overnight at 4°C with mucin fractions diluted in 4 M and 0.5 M GuHCl. Plates were incubated with a 25 mM sodium metaperiodate solution diluted in sodium acetate (NaAc) for 20 min, and 205 blocked with 50 mM Tris-HCl, 0.15 M NaCl, 90 µM CaCl₂, 4 µM EDTA, 0.01% NaN₃ and 2 % 206 bovine serum albumin, at pH 8 for 1 h. The wells were then incubated for 1 h with a biotin 207 hydrazid solution diluted 1/50 in NaAc, followed by europium labeled streptavidin diluted 208 1/1000 in DELFIA[®] Assay buffer (PerkinElmer, Waltham, MA, USA). Finally, plates were 209 incubated with DELFIA[®] enhancement solution for 5 min on a shaker. Between each step the 210 plates were washed three times with a solution containing 5 mM Tris-HCl, 0.15 M NaCl, 0.005% 211 Tween 20, and 0.01% NaN3, at pH 7.75, except for the final step where plates were washed six 212 times. Signal was measured in a Wallac 1420 VICTOR² microplate reader (PerkinElmer, 213 Waltham, MA, USA) by time-resolved fluorometry. 214

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216 MUC5AC and MUC2 enzyme-linked immunosorbent assay (ELISA)

217 Mucin fractions were diluted in 0.5 M GuHCl and coated overnight at 4°C onto 96-well plates 218 (Nunc®, Thermo Scientific). For MUC2 detection, samples were reduced with 80 µl of 2 mM 219 DTT diluted in buffer (6 M GuHCl, 5 mM EDTA, 0.1 M Tris-HCl, pH 8.0), at 37°C for 1 h. On top of the previous solution 20 µl of 5 mM IAA was added and incubated for 1 h in the dark. 220 Plates were washed three times with PBS containing 0.05% Tween-20 (PBS-T) and blocked with 221 222 1% blocking reagent for ELISA (Roche Diagnostics, Basel, Switzerland) containing 0.05% Tween 20 for 1 h, followed by incubation with the primary antibody anti-MUC5CR and anti-223 MUC2C3 (both kindly provided by G. Hansson, University of Gothenburg, Sweden) diluted 224 225 1/1000. Three more washes with PBS-T were performed before and after wells were incubated with a horse radish peroxidase (HRP) conjugated donkey anti-rabbit IgG (Jackson 226 ImmunoResearch, West Grove, PA, USA) diluted 1/10000 for 1 h. Subsequently, 100 µl of 227 228 tetramethylbenzidine substrate (Sigma-Aldrich, St. Louis, MO, USA) was added per well, and the reaction was stopped with an equivalent volume of 0.5 M H₂SO₄. Absorbance at 450 nm was 229 measured in a Wallac 1420 VICTOR² plate reader. The 45M1 antibody (Sigma-Aldrich, St. 230 Louis, MO, USA) was used to confirm the specificity of the MUC5AC signal from the isolated 231 mucins obtained with the anti-MUC5CR antibody, verifying the signal obtained with the soluble, 232 233 but not the insoluble mucins. As a result we performed a range of control analyses with and without reduction and alkylation of the mucin samples, concluding that the absence of MUC5AC 234 signal in the insoluble mucins using the 45M1 antibody was due to the destruction of the epitope 235 236 recognized by this antibody after reduction and alkylation. Thus, we are convinced the MUC5AC signal is specific, although the antibodies were designed to detect human mucins. 237

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239 Bacterial strain and culture conditions

B. hyodysenteriae strain 8dII was cultured on tryptone soy agar (TSA, Thermo Fischer Scientific,
Waltham, MA, USA) plates supplemented with 5% sheep blood (Thermo Fischer Scientific,
Waltham, MA, USA), 0.1% yeast extract (Merck, Darmstadt, Germany), 400 µg/ml
spectinomycin, 25 µg/ml colistin and 25 µg/ml vancomycin (AppliChem, Darmstadt, Germany),
at 40°C under anaerobic conditions.

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246 Mucin sample preparation and concentration estimation

Gradient fractions containing mucins were pooled together to obtain one sample for each 247 248 gradient (*i.e.* two from each pig, insoluble and soluble). Mucin concentrations in pooled samples 249 were determined by serial dilutions as well as a standard curve of a fusion protein of the mucin 250 MUC1, 16TR and IgG2a Fc (25), starting at a concentration of 20 mg/ml and using seven 1/2 251 serial dilutions in a carbohydrate detection assay described above. The mucin concentrations were calculated from the standard curve. Setting the concentration based on the glycan content 252 253 appears most appropriate as bacterial-mucin interactions largely occur via the mucin glycans (14). Although this is not an absolute measure of concentration it can be used to ensure that the 254 mucins are at the same concentration for comparative assays. Mucin concentration can also be 255 256 determined by freeze drying, however all mucins do not come into solution after freeze drying, 257 therefore this method of concentration determination can contain large errors as well as remove mucin species selectively. 258

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260 Binding of B. hyodysenteriae to pig mucins

261 White 96-well plates (Corning Life Sciences, NY, USA) were coated overnight at 4°C with 6 262 μ g/ml mucins in 0.5 M GuHCl. Wells were washed three times with PBS-T and blocked with 263 200 μ l of 5% fetal bovine serum (FBS) for 1 h. Bacteria were harvested from TSA plates 264 (above), washed in PBS, centrifuged at $2500 \times g$ for 5 min, and re-suspended in PBS with 5% FBS. One hundred microliters of a bacterial suspension diluted $to 10^8$ bacterial cells/ml were 265 added per well and the plates were shaken during incubation for 2 h at 40°C in an anaerobic 266 environment. Plates were washed three times with PBS-T and once with PBS. Subsequently, 100 267 µl of PBS was added to each well, followed by the addition of an equal volume of BacTiter-268 GloTM reagent (Promega, Madison, WI, USA). Incubation proceeded for 5 min at room 269 temperature. Relative luminescence (RLU) was measured in an Infinite® M200 microplate 270 reader (Tecan, Männedorf, Switzerland) with an integration time of 1000 ms per well. Controls 271 272 included wells without the bacteria (PBS only) in mucin coated wells, and non-mucin coated wells incubated with the bacterial suspension followed by addition of PBS and reagent. In order 273 to confirm that differences in *B. hyodysenteriae* binding to pig mucins were not due to variations 274 275 of the mucin glycan content between samples, a glycan detection assay described above was simultaneously coated and performed with the binding experiments. Only samples with a glycan 276 value of 17000-24000 Eu-counts were included to ensure that the analysis occurred within the 277 linear range of the assay, and the *B. hyodysenteriae* binding signal was normalized against the 278 glycan value for that particular coating and mucin. Results were obtained from three independent 279 experiments with five technical replicates for each mucin, and were plotted as relative 280 luminescence per glycan unit. 281

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283 Statistical analysis

Statistical analysis was performed using GraphPad Prism version 6 software (La Jolla, CA,
USA). Results are expressed as the mean ± SEM for normally distributed data, and median with
interquartile range (IQR) for data that did not follow a normal distribution (determined using the
D'Agostino-Pearson omnibus test). Data were analyzed using the Mann-Whitney, Kruskall-

Wallis or One-way ANOVA tests wherever applicable, and p values ≤ 0.05 were considered as statistically significant.

290

291 **Results**

292 Clinical signs, bacterial shedding and histology after experimental inoculation

293 Five inoculated pigs excreted *B. hyodysenteriae* in their feces and developed clinical signs of SD, including mucoid or hemorrhagic diarrhea. A milder case of diarrhea was observed in one of the 294 five pigs with clinical signs of SD. The majority of the pigs had acute dysentery at the time of 295 296 sacrifice (Table 1). Pig 4 had a longer duration of dysentery, however the clinical signs did not change during the 25 day period, and the macroscopic lesions corresponded with those of acute 297 298 dysentery. Pig 3 had recovered from clinical signs before the day it was euthanized. Fecal 299 shedding of *B. hyodysenteriae* by the inoculated pigs that developed SD started simultaneously with the onset of clinical signs and continued until the time of sacrifice. The control pigs did not 300 excrete B. hyodysenteriae in their feces and did not present any clinical signs of SD at any time 301 point during the experiment. In line with previous reports (20, 26), severe lesions were observed 302 in the colon of pigs with acute dysentery, including necrotic colitis, hyperemic mucosa and fluid 303 304 content with large amounts of mucus. Microscopically, the mucosa of pigs with SD had a 305 thickened mucus layer (Figure 1), the epithelium contained abundant goblet cells, and the colonic crypts were elongated, dilated and filled with mucus and cell debris. Inflammatory cells were 306 307 observed in the lamina propria, consisting of lymphocytes, plasma cells, and transmigrating 308 neutrophils. The control group, as well as the inoculated pigs that did not develop SD, had no 309 significant histopathological lesions.

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312 Colonic mucus disorganization during swine dysentery is accompanied by de novo expression

313 of MUC5AC and increased expression of MUC2

Fluorescence microscopy of the pig colon tissue revealed that the mucus layer in the healthy pig 314 colon, including the inoculated pigs that did not develop SD, was organized in striations parallel 315 to the mucosa, and consisted of mainly the MUC2 mucin (Figure 1, A-C), similar to the mucus 316 structure reported for the mouse colon (2, 27). In contrast, during infection with B. 317 hyodysenteriae, a massive increase in MUC2 and also de novo production of MUC5AC was 318 observed in the four inoculated pigs with severe clinical signs of dysentery (Figure 1, D-F). 319 320 MUC5AC expression was not observed by immunofluorescence in the inoculated pig with milder clinical signs of dysentery. Both MUC2 and MUC5AC mucins were produced by goblet 321 322 cells, and when MUC5AC was present in a goblet cell, it was usually present in a cell that also 323 produced MUC2 (Figure 1, D and E). In addition to the massive increase in mucus layer thickness that occurred in dysenteric pigs, the mucus organization was vastly changed by 324 infection as the striated organization was lost and instead the mucus appeared to flow in "rivers" 325 with eukaryotic cells in between, often at a 45° angle from the mucosa. 326

327

328 The antibodies used have previously been shown to detect their specific targets in humans and mice (27, 28), but no MUC2 or MUC5AC antibodies have been verified for use in pigs. To be 329 certain the stain represents MUC5AC and MUC2, we confirmed that the antibodies we used for 330 331 the immunofluorescence indeed bound to the isolated mucins in a specific manner that differed between the antibodies (Figure 2, A and B). The specificity of the MUC5AC antibody was 332 further supported by the use of a second antibody, both MUC5AC antibodies followed a tissue 333 334 distribution in the porcine stomach analogous to the distribution observed in the human and murine stomach. In addition, we designed qPCR primers specific for swine MUC2 and 335

MUC5AC, and indeed, the mRNA levels of *MUC2* and *MUC5AC* increased four-fold and more than 15-fold, respectively, in the colon tissue of pigs with clinical signs of SD compared to the control pigs (Figure 3). The expression of MUC5AC was not upregulated in the colon tissue of the inoculated pig with milder clinical signs of dysentery. The mRNA levels of *MUC1*, a mucin induced during some bacterial infections in the mouse (3, 11) did not increase in the pigs with SD compared to the controls (Figure 3).

342

343 Swine dysentery is associated with a five-fold mucin increase

344 Mucins from *B. hyodysenteriae* inoculated and control pigs were isolated from the colonic mucus 345 and analyzed in order to determine changes in their composition during infection. Mucins were 346 extracted as previously described (23) and GuHCl soluble and insoluble mucins were obtained. 347 Although insoluble mucins were ultimately solubilized by reduction in DTT they will be referred to from here on as "insoluble". During isopychnic density gradient centrifugation, molecules 348 concentrate as bands where the molecule density matches the density of the surrounding solution. 349 As mucins are highly glycosylated, and sugars have a high density, density gradient 350 centrifugation separates mucins from the less glycosylated non-mucin molecules. The initial 351 352 CsCl/4M GuHCl isopycnic density gradient procedure rendered the isolated mucins free of non-353 mucin proteins however they were contaminated with DNA (Figure 4A). Therefore a second CsCl/0.5 M GuHCl gradient was performed in all the samples ensuring removal of DNA 354 355 contamination (Figure 4B). The median mucin density of the inoculated pigs was 1.527 g/ml (IQR = 0.015). No differences in mucin density were noted between inoculated and control pigs 356 (p > 0.05, Figure 4D). Quantification of mucins based on their carbohydrate content revealed that 357 358 the pig colon mucins of both B. hyodysenteriae inoculated and control pigs were mainly insoluble, with less than 20% of the mucins being soluble in GuHCl (Figure 4C). Pigs with 359

clinical signs of SD had a five-fold higher mucin content (p < 0.05) compared to the controls (Figure 4C). The amount of mucins isolated from control pigs was similar to the mucin content of the inoculated pigs that did not develop SD (p > 0.9, Figure 4C).

363

In most density gradient samples, MUC5AC and MUC2 antibody reactivity coincided with the 364 365 glycan peak (Figure 2A), although in one sample there were differences in the MUC2 and MUC5AC curves, demonstrating that the antibodies indeed recognized different mucins (Figure 366 2B). MUC5AC was present in the GuHCl soluble and insoluble material in similar proportions 367 368 (45% and 55%, respectively, Figure 2C). Mucins from the pigs with SD contained more MUC5AC compared to the controls and to the inoculated pigs that did not develop SD (p < 0.05, 369 Figure 2C). In line with the immunofluorescence and qPCR results, the pig with mild clinical 370 371 signs of SD had the lowest level of MUC5AC antibody reactivity. Both GuHCl soluble and insoluble mucins contained MUC2, with the majority (80-90%) of MUC2 present as insoluble 372 mucin. In line with the immunofluorescence and qPCR results, the MUC2 protein level was also 373 increased in pigs with SD compared to the controls, as well as to the inoculated pigs that did not 374 develop SD (p < 0.05, Figure 2D). 375

376

Increased binding ability of B. hyodysenteriae to colonic mucins from pigs with clinical signs of swine dysentery

B. hyodysenteriae bound to colonic mucins isolated from both control and inoculated pigs. The *B. hyodysenteriae* binding pattern to mucins differed between individual pigs (insoluble mucins, overall p < 0.0001; soluble mucins, overall p < 0.0001; Figure 5A). This suggests that *B. hyodysenteriae* has an adhesin that recognizes specific glycan structure(s), as bacterial adhesins usually recognize these, and the mucin glycans differ between individuals (24). *B.* 384 hyodysenteriae binding per mucin glycan unit was higher to soluble mucins from pigs with clinical signs of SD compared to controls (p < 0.0001), and a similar trend was observed for the 385 insoluble mucins (p = 0.0595, Figure 5B). Although B. hyodysenteriae bound more to the 386 soluble mucins isolated from pig 3 compared to the other pigs with clinical signs of SD (Figure 387 5A), the overall binding difference between control and inoculated pig mucins remained 388 389 statistically significant (p = 0.0002) even after excluding pig 3 data. When taking into account the higher total mucin content isolated from pigs with clinical signs of SD than healthy pigs, the 390 total binding ability of *B. hyodysenteriae* to mucins from pigs with clinical signs of SD increased 391 392 seven-fold (p < 0.005, Figure 5C).

393

394 Discussion

395 The present study provides new insights into the composition of pig colonic mucins during health and disease as well as mucin interactions with B. hyodysenteriae. This was accomplished 396 through validation, optimization and generation of methods and tools that now can be 397 specifically applied to the swine host. We demonstrated changes in the mucus environment of the 398 swine colon during infection with *B. hyodysenteriae*, evidenced by disorganized mucus, a much 399 400 thicker mucus layer as well as five-fold higher mucin content, accompanied by de novo MUC5AC synthesis. We identified that B. hyodysenteriae bound to swine colonic mucins in a 401 manner that differed between individuals and mucin populations, and increased with infection. 402 403 As a result of these changes, the altered mucin environment provided more bacterial binding sites, increasing the overall binding ability of *B. hyodysenteriae* to colonic mucus seven-fold. 404

405

Successful isolation of mucins involves the removal of low density non-mucin proteins as well asDNA contaminants. Pure colonic mucins, soluble and insoluble in GuHCl, were obtained by two

408 isopycnic density gradient centrifugation steps in CsCl with different GuHCl molarities, as 409 previously described (29). As in human colonic mucus (30), we reported in the pig a higher 410 content of insoluble mucins compared to mucins soluble in guanidinium. Mucins are large molecules that form complex networks by connecting the mucin subunits via disulphide bonds. It 411 412 has been suggested that the higher content of insoluble mucins in the colon denotes the presence 413 of more covalent bonds, needing to be further solubilized by reduction (30). The density we observed for pig colonic mucins was higher than the density (1.38 g/ml) previously reported for 414 human colonic mucins (30). Since human and pig MUC2 are highly homologous and the 415 416 glycosylation has a similar monosaccharide composition, differences in density are likely to mainly reflect differences in the extent of glycosylation. Thus, pig colonic mucins appeared to be 417 418 more heavily glycosylated than the human counterparts. The main carbohydrates that compose 419 glycoproteins both in human and pigs are glucosamine, galactosamine, galactose, fucose and sialic acid (30-32). 420

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Besides lubricating the intestinal surface for the transit of the fecal bolus, goblet-cell secreted 422 423 mucus protects the surface epithelium from bacterial invasion. The mucus layer of healthy pigs 424 was constituted mainly of MUC2 mucin, organized in a striated fashion perpendicular to the mucosal surface, similar to the mucus composition of the mouse colon (6). During B. 425 hyodysenteriae infection, we found a loss of the striated organization, and a substantial increase 426 427 in MUC2 and *de novo* secretion of MUC5AC mucins. We recently reported dynamic changes in the mucus barrier during Citrobacter rodentium infection in mice, with structural loss and 428 decrease of the inner mucus layer at the onset and mid time points of infection (27). However, 429 430 during the clearance phase, the mucus layer thickness increased, but had a similar organization as in uninfected mice, and no Muc5ac was detected (27). The changes observed in B. 431

hyodysenteriae infected pigs are thus completely different to any of the mucus changes identified
during *C. rodentium* infection and clearance.

434

435 MUC2 and MUC5AC are both gel-forming mucins secreted by goblet cells. The MUC2 mucin is predominantly secreted in the intestine. There is evidence from Muc2 knock out mice that the 436 437 lack of Muc2 increases the susceptibility to Salmonella and C. rodentium infections (33, 34). Unlike MUC2, MUC5AC does not form part of the normal mucin repertoire in the colon. 438 Instead, it is commonly found in the normal gastric mucosa (5), airway epithelium (35) and 439 440 conjunctiva (36). MUC2 and MUC5AC mRNA levels were increased in the colon tissue of pigs with clinical signs of swine dysentery, compared to the control pigs, demonstrating that the 441 442 mucus change is regulated at the transcriptional level, in contrast to increases in mucus thickness 443 seen in C. rodentium infection without changes in mRNA levels (27). Additionally, the fact that MUC1 expression was not increased in the inoculated pigs compared to the controls further 444 supports the conclusion that B. hyodysenteriae infection has a different effect on mucin 445 regulation compared to the C. rodentium model where Muc1 is increased (27). Similar to our 446 results, expression of MUC5AC and MUC2 has been described in rabbit ileal loops inoculated 447 448 with Shigella flexneri and Shigella dysenteriae (37). In addition, Muc5ac expression is increased mice infected with the intestinal nematode Trichuris muris (38). 449 in In pigs, immunohistochemical staining with an antibody against human MUC5AC has indicated that this 450 451 mucin is increased during infection with *Salmonella* Typhimurium (39), while during infection with Trichuris suis MUC5AC mRNA levels were elevated (40). During nematode infection, 452 Muc5ac induction has a protective role in mice, decreasing nematode burden and viability (41). 453 454 Moreover, Muc5ac deficiency hampers the clearance of the parasite, increasing the susceptibility 455 to chronic infection (41). An altered mucin expression in the colon of pigs with SD was first 456 reported by Wilberts *et al.*, after immunohistochemical staining with an antibody against human 457 MUC5AC indicated its presence in pigs with acute dysentery following inoculation with B. hyodysenteriae or "B. hampsonii" (20), suggesting a common mucin response in the colon during 458 infection with these pathogens. Our results suggest that during infection, de novo secretion of 459 MUC5AC in the colon could depend on the stage of the disease, as it was not detected in the pig 460 461 sacrificed one day after the onset of clinical signs, presenting only mild diarrhea. Furthermore, the similar mucin profile between the inoculated pigs that did not develop SD and the control 462 pigs suggests that *de novo* secretion of MUC5AC in the colon depends on the ability of the 463 464 bacterium to colonize the host. Further experiments are required to determine whether the de *novo* MUC5AC secretion plays a protective role during *B. hyodysenteriae* infection in the pig. 465

466

467 Successful colonization of the host by enteric pathogens involves penetration of the mucus layer overlying the epithelium. Genomic evidence shows that B. hyodysenteriae carries genes 468 associated with potential virulence factors involved in motility, chemotaxis, and tissue injury by 469 proteases and hemolysins, that if expressed could facilitate colonization of the colon (42). Thus 470 471 far, the importance of motility and chemotaxis in B. hyodysenteriae colonization has not been 472 thoroughly demonstrated. A strong chemotactic response to pig mucins and components like 473 fucose and serine has been described (43, 44), although a decreased attraction to mucins at concentrations greater than 6 % has also been reported (45). Colonization of the gastrointestinal 474 475 tract can also be mediated by bacterial adhesion to carbohydrate structures such as blood group 476 antigens that act as receptors. For example, *H. pylori* strains that express the BabA adhesin bind to the Lewis b blood group antigen expressed in the human gastric mucosa, resulting in a blood 477 478 group and strain dependent binding (46), and the FedF adhesin expressed in F18 fimbriated E. 479 *coli* binds to glycosphingolipids isolated from intestinal epithelium of blood group A and O pigs

480 (47). In the pig intestine AO blood group antigens are expressed, with a predominance of blood 481 group A (48), and thus individual pigs carry different glycan structures in their intestines. Here 482 we showed that *B. hyodysenteriae* bound to mucins from all pigs in the study, but that the level of binding per mucin glycan unit differed between the mucin populations and with disease status. 483 In light of that other infections have previously shown to induce changes in mucin glycosylation 484 485 (49), it is likely that the differences in *B. hyodysenteriae* binding reflect differences in the pig/mucin glycan repertoire rather than differences in the mucin density or extent of 486 glycosylation. 487

488

Mucins from pigs with clinical signs of SD bound more *B. hyodysenteriae* compared to mucins 489 from the control pigs. Potentially, during infection the mucin secretion provides distinct 490 491 carbohydrate structures for *B. hyodysenteriae* binding. The mucous niche is very unstable, and pathogen binding to mucins may prevent the more intimate adherence that can occur between the 492 pathogen and, for example, glycolipids of the cell membrane. Indeed, mucin binding to the 493 human gastric pathogen H. pylori acts as a decoy and prevents prolonged adherence (13). 494 495 Furthermore, in the rhesus monkey model of *H. pylori* infection, animals with mucins that bind 496 H. pylori more effectively have a lower H. pylori density in their stomachs, indicating that mucin binding to *H. pylori* aids in removing the bacteria from the gastric niche (50). However, it is not 497 certain if these principles apply to *B. hyodysenteriae*; the massively thick disorganized mucus 498 499 layer may not be as unstable as a normal mucus layer, and there is a possibility that the protective function of the mucus changes under these conditions. B. hyodysenteriae may indeed induce 500 501 these mucus changes to create a more favorable niche instead.

In conclusion, *B. hyodysenteriae* bound to mucins from all pigs, in a manner that differed between the mucin populations and increased with SD. Together with the massive mucus induction and disorganization that occurred during infection, this demonstrates major changes in the colon mucus niche during *B. hyodysenteriae* infection.

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Pig ID (Exp. N°) [#]	Treatment group	Samples analyzed [†]	Start of clinical signs (dpi)	Days from start of clinical signs until necropsy	Clinical signs at day of necropsy	<i>Bh</i> in feces the day of necropsy	Macroscopic signs of SD at necropsy	Histological signs of SD at necropsy
1 (2)	Treatment group Inoculated with Bh		39	1	Yes		Yes	Yes
2 (2)		Yes	29	11	Yes	Yes		
3 (1)			12	28	No*			
4 (1)			15	25	Yes			
5 (2)			29	11	Yes			
6 (1)			No SD	N/A	No	No	No	No
7 (1)								
8 (1)								
9 (1)								
10-14 (1-2)		No						
A-F (1-2)	Comtral	Yes	- N/A					
G-N (1-2)	Controls	No						

687 **Table 1.** Experimental design and data of *B. hyodysenteriae* inoculated and control pigs

688 # Exp. N° states whether the pigs belonged to the 1st or 2nd infection trial; Bh = Brachyspira hyodysenteriae; dpi = days post

689 inoculation; SD = swine dysentery, N/A = not applicable; † Inoculated pigs without clinical signs of SD and control pigs were

690 randomly selected to match the number of pigs with clinical signs of SD; * Pig 3 presented clinical signs of mucoid hemorrhagic

691 diarrhea before sacrifice, and severe necrotic lesions in the colon at necropsy.

Target	Direction	Sequence (5'- 3')	Reference
MUC1	Forward	TCCGACCCGGGATGCCTACCA	This study
	Reverse	GGCTGCCCCACCGTTGCCT	This study
MUC2	Forward	CCTTGCTCTCGTGTGGAACA	This study
	Reverse	ACTTCTCCTCGGGGCTTGTTG	This study
MUC5AC	Forward	TGCGCCGTGCCACGCGGAGAT	This study
	Reverse	GCGGGGCAGGGGAAGGGGCA	This study
ACTB	Forward	CACGCCATCCTGCGTCTGGA	(22)
	Reverse	AGCACCGTGTTGGCGTAGAG	(22)
RPL4	Forward	CAAGAGTAACTACAACCTTC	(22)
	Reverse	GAACTCTACGATGAATCTTC	(22)

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Figure 1. Colon tissue sections from control and *B. hyodysenteriae* inoculated pigs stained for MUC2 and MUC5AC. Immunofluorescence of MUC2 (green) and MUC5AC (red) in colon tissue counterstained with DAPI (blue). Panels A-C show the striated organization of the mucus in the colon of control pigs along with expression of MUC2. In contrast, panels D-F show a

disorganized mucus barrier with *de novo* expression of MUC5AC and increased expression ofMUC2 in the colon of pigs with clinical signs of SD.





Figure 2. MUC5AC and MUC2 content in the colon of *B. hyodysenteriae* inoculated pigs. 703 704 (A) The peak of antibody reactivity against MUC5AC and MUC2 coincided with the glycan 705 detection peak in GuHCl soluble mucin fractions isolated from a B. hyodysenteriae inoculated 706 pig with clinical signs of SD. (B) The mucin population of one pig was more heterogeneous with distinctly different mucin peaks, demonstrating that the antibodies against MUC2 and MUC5AC 707 recognize different mucins. (C) MUC5AC and (D) MUC2 antibody reactivity against GuHCl 708 soluble and insoluble mucins isolated from *B. hyodysenteriae* inoculated pigs that developed SD 709 (**0** pig 1, **2** pig 2, **3** pig 3, **4** pig 4, and **5** pig 5), inoculated pigs that did not develop SD, and 710 control pigs. Results are expressed as the median with interquartile range. Kruskall-Wallis test 711 with Dunn's correction for multiple comparisons, * p < 0.05. 712



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Figure 3. *MUC5AC, MUC2* and *MUC1* mRNA expression of *B. hyodysenteriae* inoculated and control pigs. Normalized fold expression of *MUC5AC, MUC2*, and *MUC1* mRNA levels in the colon tissue of *B. hyodysenteriae* inoculated pigs with clinical signs of SD and controls by qPCR analysis. Expression data were normalized against *ACTB* and *RPL4* reference genes. Fold changes were calculated using $\Delta\Delta$ CT. Results are expressed as the median with interquartile range. Mann-Whitney test, * *p* < 0.05 and ** *p* < 0.005.



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Figure 4. Isolation, density and glycan content of colonic mucins from B. hyodysenteriae 723 724 inoculated and control pigs. (A) Mucin fractions were recovered from the density gradients and 725 analyzed for their glycan content. Here, a representative sample of soluble mucins isolated from a B. hyodysenteriae inoculated pig with clinical signs of SD after the first gradient in CsCl/4 M 726 GuHCl (starting density of 1.39 g/ml), shows that low density non-mucin proteins (A₂₈₀ nm) are 727 excluded from the pooled mucin fractions. Bar: pooled mucin fractions 2-7. (B) Representative 728 729 sample of soluble mucins isolated from a control pig, shows baseline separation between the 730 glycan peak and DNA after a second gradient in CsCl/0.5 M GuHCl (starting density 1.5 g/ml). 731 Bar: pooled mucin fractions 1-2. (C) Glycan content of GuHCl soluble and insoluble mucins isolated from inoculated pigs with clinical signs of SD (**0** pig 1, **2** pig 2, **3** pig 3, **4** pig 4, and 732 733 • pig 5), inoculated pigs that did not develop SD, and control pigs. The mucin content in the colon was five-fold higher in inoculated pigs with clinical signs of SD compared to the controls. 734

735 (D) Density (g/ml) of GuHCl soluble and insoluble mucins isolated from control and *B*. *hyodysenteriae* inoculated pigs (with and without clinical signs of SD) showing no differences 737 between the groups. Results are expressed as the median with interquartile range. Kruskall-738 Wallis test with Dunn's correction for multiple comparisons, * p < 0.05.





Figure 5. Binding of *B. hvodysenteriae* to colonic mucins. (A) Binding pattern of *B.* 741 hvodysenteriae to soluble and insoluble mucins isolated from control (pigs A-F) and inoculated 742 pigs with clinical signs of SD (1-5). Results are expressed as the mean \pm SEM of technical 743 replicates. One-way ANOVA, with Tukey's correction for multiple comparisons, * p < 0.05 and 744 **** p < 0.0001. (B) *B. hyodysenteriae* binding to soluble and insoluble mucins isolated from 745 746 control and pigs with SD showing higher binding to the soluble mucins isolated from pigs with 747 clinical signs of SD compared to the control group. Results are expressed as the median with interguartile range. Mann-Whitney test, **** p < 0.0001. (C) Binding ability of B. 748 hyodysenteriae to the total mucin content observed in pigs with SD (1) pig 1, 2) pig 2, 3 pig 3, 749 750 \bigcirc pig 4, and \bigcirc pig 5) and control pigs (*i.e* binding to mucin at a set concentration \times the total amount of mucin recovered from that sample). Results are expressed as the median with 751 interquartile range. Mann-Whitney test, ** p < 0.005. Data shown are representative of three 752 independent experiments. 753