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Effect of Mucine 4 and Fucosyltransferase 1 genetic variants on gut homoeostasis of growing healthy pigs

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# **ORIGINAL ARTICLE**

WILEY Animal Physiology and Animal Nutrition

# Effect of Mucine 4 and Fucosyltransferase 1 genetic variants on gut homoeostasis of growing healthy pigs 📃

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#### **Abstract**

Putative genetic markers have been associated with ETEC F4 (Mucine 4 [MUC4]; MUC4<sup>GG;CG</sup> as susceptible; MUC4<sup>CC</sup> as resistant) and F18 (Fucosyltransferate 1 [FUT1]; FUT1<sup>GG;AG</sup> as susceptible; FUT1<sup>AA</sup> as resistant) resistances respectively. In this study, 71 post-weaning pigs were followed from d0 (35 days old) to d42 (77 days of age) to investigate the effect of MUC4 or FUT1 genotypes on the mid-jejunal microbiota composition, pigs expression of genes related to inflammation (IL8, GPX2, REG3G, TFF3, CCL20 and LBPI) and glycomic binding pattern profile (Ulex europaeus agglutinin I [UEA] fucose-binding lectin and peanut as Tolinin [PNA] galactose-spe- 5 cific), and on blood plasma targeted metabolomics profile, faecal score and performance parameters of growing healthy pigs. The MUC4 and FUT1 resistant genotypes improved the pigs' growth performance and had firmed faecal score susceptible genotypes in d0-d21 period. Pigs with MUC4<sup>GG</sup> genotype had a higher jejunal expression of genes relate to immune function (CCL20 and REG3G) than MUC4<sup>CG</sup> and MUC4<sup>CC</sup> pigs (p < 0.05). MUC4<sup>CG</sup> pigs had higher expression of TFF3 (implicated in mucosal integrity) than MUC4<sup>GG</sup> and MUC4<sup>CC</sup> (p < 0.05). FUT1 influenced the alphaand beta-jejunal microbial indices. The FUT1<sup>AA</sup> group had a higher number of OTUs 6 belonging to Lactobacillus genus, while FUT1<sup>GG</sup> group had a higher number of OTUs belonging to Veillonella genus. MUC4<sup>CC</sup> pigs had lower scores for UEA on brush borders and goblet cells in villi than MUC4<sup>GG</sup> (p < 0.05). FUT1<sup>AA</sup> pigs had lower UEA positivity and higher PNA positivity on brush borders and goblet cells than FUT1AG and  $FUT1^{GG}$  (p < 0.05). Both FUT1 and MUC4 influenced the metabolic profile of healthy pigs. Results highlight the role of MUC4 and FUT1 on pig intestinal homoeostasis and improved the knowledge regarding the potential interaction between host genetics, gut microbiota composition and host metabolism in a healthy status.

#### KEYWORDS

16S rRNA, fucose, FUT1, intestinal mucosa, MUC4



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# 1 | INTRODUCTION

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In pigs, the post-weaning period is frequently characterized by suboptimal growth and a high prevalence of intestinal disorders, mainly due to colibacillosis infection (Schokker et al., 2015), that impact on the pig morbidity and mortality and increase the use of antibiotics (Gresse et al., 2017). Preserving the gut homoeostasis by maintaining a favourable intestinal bacteria balance and preserving the functionality of the mucosa to ensure the digestion and absorption of nutrients and the immunological functions are crucial to sustain the gut health (Wells et al., 2017).

Several genetic markers have been associated with pigs postweaning diarrhoea (PWD) due to enterotoxigenic Escherichia coli (ETEC) F4 and F18. The single nucleotide polymorphism (SNP) 8227C>G located on Mucine 4 (MUC4) gene has been associated with the expression of the F4ac receptor in the jejunum brush border and with the susceptibility to ETEC F4. Piglets with MUC4<sup>CG/</sup> GG genotypes result as susceptible and express F4 receptor, while piglets with MUC4<sup>CC</sup> genotype are most associated with the resistant phenotype (Jørgensen et al., 2003). The SNP located on Fucosyltransferase 1 (FUT1) gene has been associated with piglet susceptibility to ETEC F18ab/ac. Piglets with FUT1<sup>AG/GG</sup> genotypes result as susceptible; piglets with FUT1<sup>AA</sup> genotype results as resistants (Meijerink et al., 2000). FUT1 encodes for the FUT1 enzyme which catalyses the addition of terminal alpha (1,2) fucose residues on the carbohydrates expressed on the surface of epithelial cells and 7 in mucosal secretions (Henry et al., 1996) that has been highlighted as a putative receptor for ETEC F18 adhesion in the small intestine. For this reason, piglets with different FUT1 genotypes have been associated with a different phenotype for ETEC F18 receptors in the jejunum brush border and with their relative susceptibility to ETEC F18 (Coddens et al., 2007).

Both host genetic background and intestinal bacteria can influence the intestinal gut homoeostasis, and the overall host metabolism and health status (McKnite et al., 2012; Sommer & Backhed, 2013). Studies suggested that MUC4 and FUT1 genotypes are not only associated with piglets susceptibility to PWD but can also influence the intestinal homoeostasis. For instance, healthy piglets with different MUC4 and FUT1 genotypes are known to differ for their intestinal microbial profile (MUC4, Messori, Trevisi, Simongiovanni, Priori, & Bosi, 2013; FUT1, Poulsen et al., 2018) and for their intestinal protein glycosylation (Hesselager, Everest-Dass, Thaysen-Andersen, Bendixen, & Packer, 2016). Furthermore, the intestinal mucosa of piglets with the susceptible genotype (MUC4<sup>CG/GG</sup>) had an up-regulation of genes related to antimicrobial peptide and immune function such as lipopolysaccharide binding protein (LBPI) and regenerating islet-derived 3 gamma (REG3G) compared to pigs with a resistant genotype (Trevisi et al., 2012).

Those differences in the intestinal characteristics can, therefore, concur to determine differences in piglets' metabolism (Poulsen et al., 2018) and in piglet nutritional requirement (Trevisi et al., 2012) and result in different growth performance as already reported by

Fontanesi et al.. (2012) for MUC4 and by Bao et al. (2011) for FUT1 in adult pigs.

However, the effects of *MUC4* and *FUT1* genetics variances in influencing the intestinal homoeostasis and microbial profile in healthy piglets are still poorly investigated.

We hypotheses that MUC4 and FUT1 genetic variances can influence the glycomic pattern and the microbial profile of the small intestine giving a different stimulation of the mucosal immune response, resulting in the modulation of piglets' metabolism and growth performance. The aim of the present study was to evaluate the effect of MUC4 and FUT1 genetic variances on the growth performance, physiological status and the intestinal homoeostasis of growing healthy pigs.

#### 2 | MATERIAL AND METHODS

The procedures complied with the Italian law pertaining to experimental animals and were approved by the Ethic-Scientific Committee for Experiments on Animals of the University of Bologna and the Italian Ministry of Health by the approval number 801/2015-PR.

# 2.1 | Animal and sampling

One week post-weaning (35 days of age - d0), seventy-one piglets were randomly selected from 10 litters based on their individual body weight (7,063 ± 936 g average body weight), moved to the experimental facility of the University of Bologna and located in individual pens with a mesh floor for 6 weeks (d42). The study was carried out in three consecutive batches composed by 24, 24 and 23 pigs respectively. Pigs were kept at a controlled temperature (28°C at d0 to 25°C at d42) and heated additionally by infrared lamps for the first 7 days. Animals were fed two consecutive diets: a pre-starter diet from d0 to d21 and a starter diet from d22 to d42 (Table 1). Feed intake was daily recorded, and piglets were individually weighted every week until the end of the trail. To evaluate the pigs' health status, the individual faecal score was daily recorded by visual appraisal using a five-point scoring system (1-5): 1: hard, 5: watery faeces, where a faecal score >3 was considered as diarrhoea. Weekly faecal score was then calculated individually. From each pig, bristles were collected for the genotyping analysis. On d42, all animals were anaesthetized and sacrificed via an intracardiac injection of Tanax (0.5 ml/kg body weight). The jejunum was isolated at 50% of the small intestine length. Jejunum content was collected and mixed with the mucous layer obtained from a gentle scraping of the jejunum mucosa. The sample was snap-frozen in liquid nitrogen and stored at -20 °C for microbiota analysis. From the same emptied jejunal site, the mucosa was gently scraped, snap-frozen in liquid nitrogen and preserved at -80°C for gene expression analyses.

From a sub-group of 24 pigs, selected to be balanced for FUT1 and 8 MUC4 genotypes, an additional sample of mid-jejunum tissue was collected, washed with 0.01 M phosphate-buffered saline (PBS), pinned tightly to balsa wood, fixed in 10% buffered formalin for

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**TABLE 1** Ingredients and composition of piglet pre-starter diet (from d0 to d21) and starter diet (from d22 to d42) expressed on a dry matter basis

| n y matter basis              |                  |              |
|-------------------------------|------------------|--------------|
|                               | Pre-starter diet | Starter diet |
| Ingredients (%)               |                  |              |
| Barley                        | 25.00            | -            |
| Corn                          | -                | 56.00        |
| Wheat                         | 21.00            |              |
| Wheat bran                    | -                | 18.00        |
| Cereal flakes                 | 12.00            | -            |
| Milk whey                     | 11.00            | -            |
| Soya bean meal                | 4.80             | 20.00        |
| Soya bean protein concentrate | 11.00            | -            |
| Cooked soy grains             | 8.00             | -            |
| Corn starch                   | 1.60             | 1.50         |
| Sunflower oil                 | 1.00             | 0.50         |
| L-Lysine HCl                  | 0.66             | 0.60         |
| L-Threonine                   | 0.30             | 0.25         |
| L-Tryptophan                  | 0.08             | 0.07         |
| DL-Methionine                 | 0.30             | 0.24         |
| <sub>L</sub> -Valine          | 0.15             | 0.16         |
| Salt                          | 0.45             | 0.45         |
| Dicalcium phosphate           | 0.86             | 0.73         |
| Calcium carbonate             | 1.30             | 1.00         |
| Vitamin and mineral premix    | 0.50             | 0.50         |
| Composition (as fed)          |                  |              |
| Net energy (MJ/kg)            | 10.36            | 9.6          |
| Crude protein (g/kg)          | 18.5             | 17.3         |
| Digestible lysine (g/kg)      | 1.31             | 1.18         |
| Calcium (g/kg)                | 0.98             | 0.65         |
| Digestible phosphorus (g/kg)  | 0.38             | 0.22         |

24 hr at room temperature (RT) and dehydrated in a graded series of ethanol and embedded in paraffin for immunohistochemistry analysis. A blood sample from the same sub-group was obtained by venipuncture of vena cava on a collection tube with K3 EDTA (Vacutest Kima Srl, Padova, Italy) before the animal sacrifice. Blood samples were centrifuged at 3,000 g for 10 min at 4°C to obtain the plasma. Plasma samples were stored at -80°C until further targeted metabolomics profile analysis.

# 2.2 | Animal genotyping

Genomic DNA of each pig was extracted from bristles. The bristle bulbs were incubated in Proteinase K solution (10 mg/ml of proteinase K in buffer [20 mM Tris–HCl (pH 8.4), 50 mM KCl]) for 2 hr at 50°C; then, the proteinase was inactivated at 95°C. Samples were stored at –20°C. Genotyping of the MUC4 g.8227C>G, FUT1 g.307 G>A, was carried out by Restriction Fragment Length Polymorphism

PCR (PCR-RFLP) using specific primers, annealing temperature, condition enzymes reported by Jørgensen et al. (2003) and Meijerink et al. (2000).

# 2.3 | Gene expression

The total RNA was isolated from scraped mid-jejunum mucosa using the TRIzol reagent (Life Technologies, Carlsbad, CA, USA) extraction. Quantification and quality control of all RNA samples were assessed using the ND 1000 Spectrophotometer (NanoDrop Technologies Inc., Wilmington, DE, USA) with an optimal ratio 260/280 nm between 1.7 and 2.2 and visualized on 1.5% of agarose gel. cDNA was synthesized using 1,000 ng of RNA as template (ImProm-II Reverse Transcription System Promega Corporation, Milan, Italy). The semiquantitative PCR (qPCR) of interleukin-8 (IL8), glutathione peroxidase (GPX2), regenerating islet-derived 3 gamma (REG3G), trefoil factor 3 (TFF3), C-C motif chemokine ligand 2 (CCL20), ST3 beta-galactoside alpha-2,3-sialyltransferase 1 (ST3GAL), lipopolysaccharide binding protein (LBPI) were performed in a Light Cycler instrument (Roche, Mannheim, Germany). The genes related to immune function (IL8, CCL20 and REG3G), antimicrobial activity (LBPI) and inflammatory response (GPX2, TFF3) were selected because highlighted in several studies as effective markers able to disclose alterations of tle the mucosa homoeostasis, especially in ETEC-infected pigs (Bosi et al., 2004; Sargeant et al., 2010; Trevisi, Latorre et al., 2017; Trevisi 9 10 et al., 2018).

Primers' sequence, amplified fragments' length and qPCR conditions are reported in Luise et al. (2017). The amplification was carried out in 10  $\mu$ l overall volume containing 2  $\mu$ l of cDNA, 8 pmol of each primer and 5  $\mu$ l of SYBRs Premix Ex TaqTM II (Perfect Real Time; Takara Bio Inc., Shiga, Japan). The relative quantification of gene expressions was calculated using standard curve methods and normalized to a housekeeping gene hydroxymethylbilane synthase (HMBS).

#### 2.4 | Microbiota Analysis

Total bacterial DNA was extracted from all samples using QIAamp Stool Mini Kit (Qiagen, Hilden, Germany). DNA concentration and its purity (absorbance ratio 260/280 and 260/230) were measured by ND 1000 Spectrophotometer (NanoDrop Technologies Inc., Wilmington, DE, USA) and visualized on 1.5% of agarose gel. All samples had absorbance ratios 260/280 higher than 1.8 and 260/230 ratios lower than 2. In addition, a PCR for lactic acid bacteria (LAB) was performed following the protocol reported by Walter et al. (2001), to further confirm that bacterial DNA was extracted. The library formation and sequencing of the 16S rRNA gene were performed with MiSeq® Reagent Kit V3-V4 on MiSeq-Illumina® platform. Generated sequences (approximately ~460 bp) were analysed using subsampled open-reference OTU strategy with default settings in QIIME (v1.9.1) (Caporaso et al., 2010). The reads of the 16S rRNA gene were paired-end and demultiplexed. Subsampled open-reference OTU-picking was performed

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using UCLUST with 97% sequence similarity. Sequences were chimera checked using Chimera Slayer with default settings and taxonomy assigned against the Greengenes database V13\_8 using the UCLUST method with a 90% confidence threshold. The singletons and OTUs with relative abundance across all samples below 0.005% were removed.

#### 2.5 | Immunohistochemistry analysis

Transvers µm thick) sections were obtained from jejunum and analysed Tot Ulex europaeus agglutinin I (UEA; Vector Laboratories, Peterborough, UK; catalogue n. B-1065- fucose-specific) and peanut agglutinin (PNA; Vector Laboratories, Peterborough, UK; catalogue n. B-1075- galactose-specific) following the protocol described by Priori et al. (2016). For each slide, 20 villi and 20 crypts were randomly observed. The positivity for UEA and PNA was assessed by the same operator in correspondence with the surface of the villi (microvilli brush border) and for labelled cells frequency in the villi 11 and crypts. The intensity of immunoreactivity in the surface of villi was scored as follows: 0 = absence of immunoreactivity; 1 = positivity of immunoreactivity distributed throughout the surface of the villi; and 3 = very marked positivity of immunoreactivity and great thickness. The number of immunoreactive (IR) goblet cells in the villi and crypts was evaluated and scored as follows: 1 = absent/rare IR cells; 2 = few IR cells; 3 = some IR cells; and 4 = many IR cells.

### 2.6 | Plasma targeted metabolomics analysis

Plasma metabolites were measured using the Biocrates AbsoluteIDQTM p180 Kit (Biocrates Life Science AG, Innsbruck, Austria) that allows to quantify a panel of 188 compounds, including 40 acylcarnitines, 21 amino acids, 21 biogenic amines, 90 glycerophospholipids (14 lysophosphatidylcholines [lysoPC] and 76 phosphatidylcholines [PC]), 15 sphingolipids and hexoses (sum of hexoses, including glucose). The kit was processed according to manufacturer instructions and analysed on the Serie 200 high-pressure liquid chromatography system by Perkin Elmer (Waltham, MA) coupled with the API 4000 QTRAP by AB Sciex (Foster City, CA, USA). Instrumental data were acquired and processed by Analyst 1.6.3, whereas data quantitation and validation were performed by MetIDQ-5.5.4-DB100-Boron-2623 software. Results were exported in micromolar unit (μM).

#### 2.7 | Statistical analysis

Except for microbiota and metabolomics data, all data were analysed on SAS software (version 3.4 SAS Institute), using the GLM procedure including *MUC4* and *FUT1* genotypes, litter within the batch and batch as fixed factors. Sex was included in the model when significant. A restricted model including only significant factors was then carried out.

Biostatistics on OTUs table was performed using the vegan and phyloseq packages in R software (v.3.3.0). The richness and

alpha-diversity indices (Chao1 and Shannon indices) were calculated on raw data while beta-diversity ordination and differential abundance analysis were carried out after rarefaction correction. Alpha-diversity index values were compared with multivariate ANOVA testing litters and the genotypes analysed as explanatory variables. The genotype differences in beta diversity were explored using permutational MANOVA (Adonis procedure) on Bray-Curtis distance matrix, and differences were tested by pairwise Wilcoxon signed-rank post hoc test. In order to identify the discriminant OTUs belonging to each variant of FUT1 (the sole significant gene), the sparse partial least squares discriminant analysis (sPLS-DA) was carried out using the MixOmics (mixOmics\_6.1.3) package. Microbial data were previously normalized using the total sum scaling normalization coupled with the centred log-ratio (CLR) transformation as recommended by MixOmics. The optimal number of components and the optimal number of selected variables for each component included in the sPLS-DA model were selected based on the averaged balanced classification error rate with centroids distance over 100 repeats of a 10-fold cross-validation of a sPLS-DA model. To validate the results, the stability frequency scores of the selected OTUs were calculated ("perf" function) with 10-fold cross-validation and 100 repetitions. The plotVar function was then used to identify the structure of the correlation between OTUs matrix and FUT1 genotypes. OTUs showing a correlation with FUT1 > 0.5 and a stability ≥65% were considered discriminative.

For targeted metabolomics analysis, data were imported into the MetaboAnalyst 2.0 software (http://www.metaboanalyst). Metabolites with more than 20% of missing value were removed, and the remaining missing values were calculated using the Kpp algorithm of K-Nearest Neighbours approach. Data were then normalized by each metabolite median, log-transformed and mean centred. An ANOVA model for each genotype was then carried out for all the metabolites, and *P* values were corrected for false discovery rate. Furthermore, the sPLS-DA method was applied to describe the *FUT1* and *MUC4* genotype variant influence the targeted metabolome profile and to obtain the most discriminate metabolites among the different genotypes. *p* value <0.05 was considered as significant.

#### 3 | RESULTS

#### 3.1 | Animal genotypes

For MUC4, 38 pigs had MUC4<sup>CC</sup> genotype, 26 pigs had the MUC4<sup>CG</sup> genotype and seven had the MUC4<sup>GG</sup> genotype where the MUC4<sup>GG</sup> is the susceptible genotype for ETEC F4. For FUT1, 33 pigs FUT1<sup>GG</sup>, 31 pigs FUT1<sup>AG</sup> and seven pigs FUT1<sup>AA</sup> were observed, where the FUT1<sup>AA</sup> is the resistant genotype for ETEC F18. A total of 28 pigs had the genotype indicating the susceptibility to both ETECs (three had MUC4<sup>GG</sup>, FUT1<sup>GG</sup>; three had MUC4<sup>GG</sup>, FUT1<sup>AG</sup>; 11 had MUC4<sup>CG</sup>, FUT1<sup>GG</sup>; and 11 had MUC4<sup>CG</sup>, FUT1<sup>AG</sup>). Four pigs had the susceptible genotype for ETEC F4 and the resistant one for the F18 (two had MUC4<sup>CG</sup>, FUT1<sup>AA</sup> and two had MUC4<sup>CG</sup>, FUT1<sup>AA</sup>). A total of 32 pigs had the resistant genotype for the ETEC F4 and the susceptible



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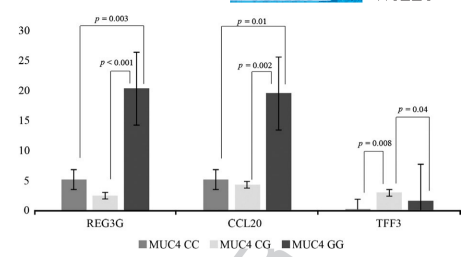
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FIGURE 1 Effect of MUC4 genotypes on the expression of REG3G, CCL20 and TFF3 in the mid-jejunal mucosa of healthy young pigs. Gene expression of REG3G, CCL20 and TFF3 and P-values of GLM model for the comparisons between MUC4<sup>CC</sup>, MUC4<sup>CG</sup> and MUC4<sup>GG</sup> pigs. Data are expressed as the ratio of mRNA copy number of target gene and mRNA copy 13 number of HMBS, multiplied by 10



one for the ETEC F18 (13 pigs had MUC4<sup>CC</sup>, FUT1<sup>AG</sup> and 19 pigs had MUC4<sup>CC</sup>, FUT1<sup>GG</sup>). One pig had the genotype indicating the resistance against both ETECs, showing the variants MUC4<sup>CC</sup> and FUT1<sup>AA</sup>.

# 3.2 | Monitoring of pigs' health status and growth performance

The pigs remained healthy throughout the experiment. The average weekly faecal score was lower than 3 (cut-off value for diarrhoea). Results showed that MUC4<sup>CC</sup> genotype had a lower (harder) faecal score than MUC4<sup>CG</sup> and MUC4<sup>GG</sup> genotypes during the second week of the study (p < 0.05). The MUC4<sup>CG</sup> tended to have higher (softer) faecal score than  $MUC4^{GG}$  during the fourth and fifth weeks (p < 0.1). The FUT1<sup>AA</sup> genotype had a faecal score lower than FUT1<sup>GG</sup> genotype during the first week and tended to have a faecal score lower during the third week. During the third week, the FUT1<sup>AA</sup> genotype had a faecal score lower than  $FUT1^{AG}$  genotype (p < 0.05; Supporting

# 12 Information Table S1).

MUC4<sup>CC</sup> genotype had a BW higher than MUC4<sup>GG</sup> genotype at d21 and d42 (p < 0.05). MUC4<sup>CC</sup> genotype had an ADG higher than MUC4<sup>GG</sup> and MUC4<sup>GG</sup> genotypes in the period from d0 to d21 (p < 0.05) and higher than  $MUC4^{GG}$  genotype in the period from d0 to d42 (p < 0.05). Feed intake was higher in MUC4<sup>CC</sup> genotype than in  $MUC4^{GG}$  genotype from d0 to d21 (p < 0.05). F:G ratio was lower in MUC4<sup>CC</sup> genotype than MUC4<sup>GG</sup> and MUC4<sup>GG</sup> genotypes in the period from d0 to d21 (p < 0.05). FUT1<sup>AA</sup> genotype had a BW at d0 lower than FUT1<sup>GG</sup> and an ADG higher than FUT1<sup>AG</sup> and FUT1<sup>GG</sup> in the period from d0 to d21 (p < 0.05; Supporting Information Table S2).

## 3.3 | Jejunum gene expression

Results showed that MUC4<sup>GG</sup> genotype had a higher expression of CCL20 and REG3G genes than MUC4<sup>CC</sup> and MUC4<sup>CC</sup> genotypes (p < 0.05). For TFF3 expression, the MUC4<sup>CG</sup> genotype had a higher expression than  $MUC4^{GG}$  and  $MUC4^{CC}$  (p < 0.05; Figure 1). FUT1 genotype did not influence the jejunum expression of the selected genes.

# Microbiota composition of mid-jejunum

Six samples were excluded from the microbiota analysis due to a low quality of sequencing data. The 3,472,457 reads obtained after quality and abundance filtering were assigned to 483 taxa and nine bacterial phyla. Firmicutes was the most represented phylum (80%), followed by Actinobacteria (8.9%) and Proteobacteria (7.9%). A total of 137 genera were identified in data aggregated at the genus level. Lactobacillus genus (39.2%) was the most abundant, followed by Streptococcus (19.8%). MUC4 polymorphism did not influence the alpha- (Chao1 and Shannon indices) and beta-diversity (Bray-Curtis distance) indices.

FUT1 polymorphism did not influence the Chao1 index, while the Shannon index differed for FUT1. The FUT1<sup>AG</sup> group (3.02  $\pm$  0.72) had a Shannon value higher than FUT1<sup>GG</sup> (2.76 ± 0.76) and FUT1<sup>AA</sup>  $(2.26 \pm 1.01)$  groups (p = 0.01). A significant effect was observed for FUT1 polymorphism (p = 0.004) on Bray-Curtis distance matrix. The pairwise adonis procedure showed a difference between the FUT1<sup>AA</sup> genotype and the other two genotypes (FUT1<sup>AA</sup> vs. FUT1<sup>AG</sup>, p = 0.01;  $FUT1^{AA}$  vs.  $FUT1^{GG}$ , p = 0.02). Results of the sPLS-DA for FUT1 were reported in Table 2. The FUT1<sup>AA</sup> group was discriminated by a higher number of OTUs belonging to Lactobacillus genus, mainly belonging to Lactobacillus reuteri species, while the FUT1<sup>GG</sup> group was discriminated by a higher number of OTUs belonging to Veillonella genus.

#### 3.5 | Plasma targeted metabolome profile

The MS/MS targeted analysis performed on plasma samples of the sub-selected group of pigs provided results for 134 metabolites (Supporting Information Table S2). ANOVA indicated that carnitine (C0), O-acyl-O-acyl phosphatidylcholine (PC\_aa) C36:1 and asymmetric dimethylarginine (ADMA) differently responded to the MUC4 genotypes and spermidine to the FUT1 genotypes. Higher level of CO was observed for  $MUC4^{CC}$  genotype (5.92  $\mu$ M) than  $MUC4^{CG}$  (4.53  $\mu$ M) and  $MUC4^{GG}$  (4.50  $\mu$ M); a lower amount of PC aa C36:1 was observed in MUC4  $^{CC}$  (32.90  $\mu$ M) than MUC4  $^{CG}$  (36.80  $\mu$ M) and

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| FUT1 <sup>a</sup> | Family           | Genus         | Species | Value.var <sup>b</sup> | Freq <sup>c</sup> |
|-------------------|------------------|---------------|---------|------------------------|-------------------|
| AA                | Lactobacillaceae | Lactobacillus | NA      | 0.19                   | 0.67              |
| AA                | Lactobacillaceae | Lactobacillus | NA      | 0.23                   | 0.77              |
| AA                | Lactobacillaceae | Lactobacillus | NA      | 0.20                   | 0.71              |
| AA                | Lactobacillaceae | Lactobacillus | NA      | 0.18                   | 0.70              |
| AA                | Lactobacillaceae | Lactobacillus | NA      | 0.10                   | 0.65              |
| AA                | Lactobacillaceae | Lactobacillus | NA      | 0.28                   | 0.81              |
| AA                | Lactobacillaceae | Lactobacillus | NA      | 0.22                   | 0.75              |
| AA                | Lactobacillaceae | Lactobacillus | mucosae | 0.23                   | 0.72              |
| AA                | Lactobacillaceae | Lactobacillus | reuteri | 0.30                   | 0.79              |
| AA                | Lactobacillaceae | Lactobacillus | reuteri | 0.26                   | 0.75              |
| AA                | Lactobacillaceae | Lactobacillus | reuteri | 0.39                   | 0.80              |
| AA                | Lactobacillaceae | Lactobacillus | reuteri | 0.24                   | 0.74              |
| AA                | Lactobacillaceae | Lactobacillus | reuteri | 0.27                   | 0.78              |
| AA                | Lactobacillaceae | Lactobacillus | reuteri | 0.22                   | 0.75              |
| AA                | Lactobacillaceae | Lactobacillus | reuteri | 0.25                   | 0.77              |
| GG                | Veillonellaceae  | NA            | NA      | 0.51                   | 0.65              |
| GG                | Veillonellaceae  | Veillonella   | dispar  | 0.48                   | 0.68              |

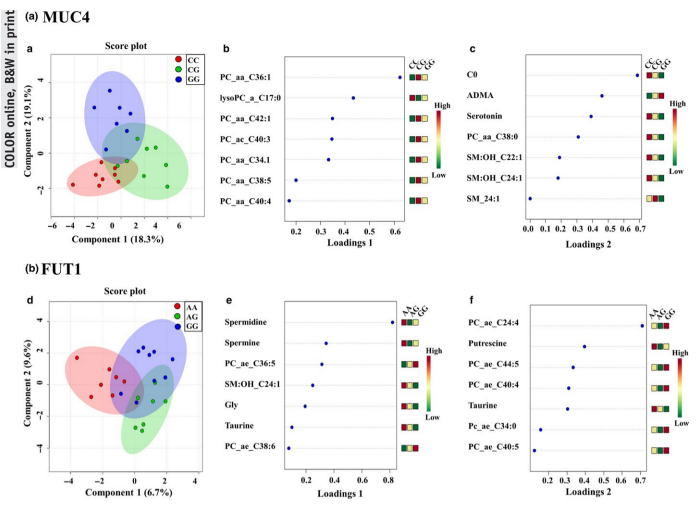
TABLE 2 Significant discriminant OTUs for FUT1 polymorphism obtained using the sparse partial least squares discriminant analysis methods coupled with 10-fold cross-validation and 100 repetitions

<sup>a</sup>FUT1, 1-fucosyltransferase gene. Genotypes had the following distribution: FUT1<sup>AA</sup> = 5; FUT1<sup>AG</sup> = 27; FUT1<sup>GG</sup> = 33. <sup>b</sup>Value.var expresses the variance explained by the single OTU. <sup>c</sup>Freq expresses the frequencies by which the OTUs were chosen among the 100 repetitions of the cross-validation.

MUC4<sup>GG</sup> variants (36.36 μM); and a higher level of ADMA was observed for MUC4  $^{GG}$  (2.42  $\mu$ M) than MUC4  $^{CC}$  and MUC4  $^{CG}$ (1.68  $\mu$ M; p < 0.05). A higher level of spermidine was observed in the FUT1  $^{AA}$  genotype (0.53  $\mu M)$  than the FUT1  $^{AG}$  (0.24  $\mu M)$ 14 and  $FUT1^{GG}$  variants (0.29  $\mu$ M; p < 0.05). The multivariate analysis showed that metabolomics profile was partially affected by MUC4 and FUT1 genotypes variants as shown by the sPLS-DA score and loading plot (Figure 2a,b). The score plot showed that MUC4<sup>CC</sup> and MUC4<sup>GG</sup> clusters were mainly distinguished by the PC2 (9.1% explained variance); MUC4<sup>CC</sup> group had a higher level of CO, serotonin, PC\_aa\_38:0, sphingomyelin (SM)OH\_C22:1, SM\_OH\_C24:1 than MUC4GG. The PC1 partially separated the MUC4<sup>CG</sup> cluster from MUC4<sup>CC</sup> and MUC4<sup>GG</sup> clusters. The MUC4<sup>CG</sup> group was characterized by a higher value of PC\_aa\_C42:1, PC\_aa\_C40:3, PC\_aa\_34:1, PC\_aa\_C38:5, and PC\_aa\_C40:4 than MUC4<sup>CC</sup> and MUC4<sup>GG</sup> groups. MUC4GG group had higher ADMA and PC\_aa\_C36:1 than the other two groups. For FUT1, the sPLS-DA score plot showed that clusters representing FUT1AG and FUT1GG clustered together along PC1 (6.7% of explained variance) while the cluster representing FUT1<sup>AA</sup> was partially separated. The loading score plot for FUT1 showed that levels of spermidine, spermine, SM\_OH\_C24:1, glycine, putrescine and taurine were higher in the FUT1<sup>AA</sup> group than FUT1<sup>AG</sup> and FUT1<sup>GG</sup> groups, while the FUT1<sup>GG</sup> group was characterized by higher level of PC\_ae\_C36:5, PC\_ae\_C42:4, PC\_ae\_C44:5, and PC\_ae\_C40:4 than FUT1<sup>AA</sup> and FUT1<sup>AG</sup> groups.

# **Immunohistochemistry**

Ulex europaeus, agglutinin I and PNA immunoreactivities were observed on the surface of the jejunum villi, while IR goblet cells were observed both in villi and in crypts and scored as shown in Figure 3. Immunoreactivity in the villi showed up as a layer that followed the entire profile of the villus. The IR goblet cells presented a typical columnar/pear shape morphology containing mucin granules intensely marked and were distributed along the axis of the villus and/or aggregated in the crypts. Opposite intensities for UEA and PNA were observed in the same brush borders. The degree of intensity seen in the goblet cells with PNA was averagely lower than with UEA. The effect of MUC4 and FUT1 genotypes on the average scores for the UEA or PNA immunoreactivity in the jejunal brush borders and IR goblet cells is shown in Table 3. The MUC4<sup>CC</sup> pigs had a lower score for UEA in brush borders and goblet cells in villi than MUC4<sup>GG</sup> (p < 0.05) and  $MUC4^{CG}$  pigs (marginally, p = 0.081 for brush borders; p < 0.05 for goblet cells). No effect of MUC4 on the UEA-IR goblet cells in crypt and of PNA for brush borders and goblet cells in villi and crypts was found. The FUT1<sup>AA</sup> genotype had lower UEA immunoreactivity score in brush border than FUT1<sup>AG</sup> and FUT1<sup>GG</sup> genotypes (p < 0.001). UEA-IR goblet cells in the villi and crypts were less in the  $FUT1^{AA}$  genotype than in  $FUT1^{AG}$  genotype and in  $FUT1^{GG}$  genotype (p < 0.05). FUT1<sup>AG</sup> had less UEA-IR goblet cells in crypts (p < 0.05), and a trend of significant lower UEA immunoreactivity score on the brush border (p < 0.1) than  $FUT1^{GG}$ . No difference was observed for UEA-IR goblet cells in crypts. For PNA, the FUT1<sup>AA</sup> genotype had



**FIGURE 2** Score plots and loading plots of sparse partial least squares discriminant analysis (sPLS-DA) on serum target metabolomics profile of young healthy pigs according to MUC4 (A) and FUT1 (B) genotypes. The (a and d) show the individual score plot for MUC4 and FUT1 variances respectively. The (b and c) represent the loading plots showing the metabolites that contribute toward the separation of sPLS-DA scores between MUC4 variances. The (d and e) represent the loading plots showing the metabolites that contribute toward the separation of sPLS-DA scores between FUT1 variances

higher PNA immunoreactivity score in brush border than  $FUT1^{AG}$  (marginally, p = 0.062) and  $FUT1^{GG}$  (p = 0.006) genotypes. PNA-IR goblet cells in the villi and crypts were more abundant in the  $FUT1^{AA}$  genotype than in  $FUT1^{AG}$  genotype and in  $FUT1^{AG}$  genotype (p < 0.05). No difference between  $FUT1^{AG}$  and  $FUT1^{GG}$  for PNA immunoreactivity in bush border and goblet cells was observed.

# 4 | DISCUSSION

The present study investigated the effect of *MUC4* or *FUT1* genotypes on gut homoeostasis of pigs by evaluating their effect on the host response in the absence of specific infections during the postweaning phase.

Our study highlighted that the *MUC4* SNP, which is associated with piglet susceptibility for ETEC F4ac (Jørgensen et al., 2003), can influence piglet growth performance and the faecal score during the first weeks post-weaning. The pigs had no diarrhoea and remained

apparently healthy during the test; nevertheless, our results indicate that the pigs with the resistant genotype (MUC4<sup>CC</sup>) had firmer faeces than the pigs with the susceptible genotypes. In addition, our study indicates that improved growth performance and improved feed conversion can be observed in the post-weaning pigs with the resistant genotype, in accord with the study of Trevisi et al. (2009) in which post-weaning piglets susceptible to ETEC F4 and fed a stan-18 dard nutritional diet had a lower ADG and higher feed intake. This could be explained by the different intestinal homoeostasis of pigs differ for the genotypes. This hypothesis is supported by the upregulation of genes related to inflammation and immune response such as REG3G, CCL20 and TFF3 observed in piglets with the susceptible genotypes in our study. These results are consistent with the findings of Trevisi, Latorre et al. (2017) for ETEC-infected pigs. Particularly, REG3G protein is mainly expressed in the intestinal crypts; it is associated with the host C-type lectins and with antimicrobial activity, and indeed, REG3G expression increases in infected pigs (Soler, Miller, Nöbauer, Carpentier, & Niewold, 2015; Trevisi et

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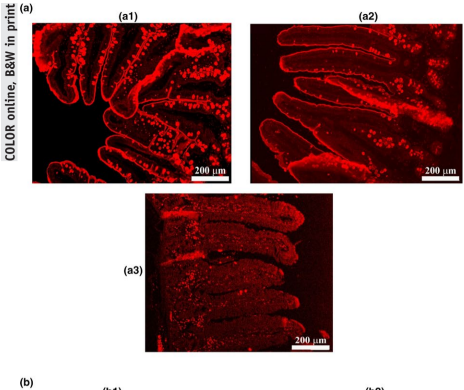
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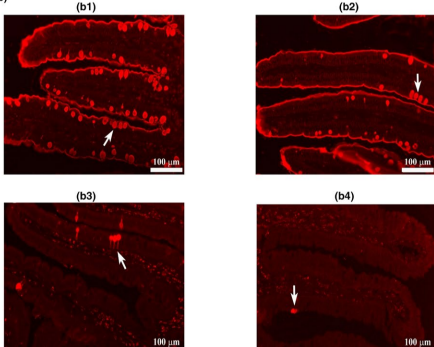


FIGURE 3 Images referred to the immunofluorescence method obtained using Ulex europaeus agglutinin I (UEA, fucose-binding) and peanut agglutinin (PNA; galactose-binding) in the pig jejunum sections for brush border (in 3.A) and for the distribution of immunoreactive (IR) goblet cells in the villi (in 3.B). The image A.1 shows the UEA high immunoreactivity of brush border in correspondence of the mucosal surface with score 2. In the villages of image A.2 16 represents the mucosal surface with score 1 (less intense and thick than previous image A.1). The image A.3 represents the mucosal surface with score 0 devoid of any immunofluorescence at the brush border. The image B.1 (UEA marker) shows the relevant number of IR goblet cells that scored 4: Many IR goblet cells with round and/or pear shape were observed along the axis of the villus. The image B2 image shows the mucosal surface of samples scoring 3 (PNA marker): Some isolated or grouped IR goblet cells were observed (arrow). The image B.3 shows the mucosal surface of samples scoring 2: The number of immunofluorescent cells (slag) is reduced (UEA marker). Rare IR cells (score 1) are shown in the image B.4 (PNA 17 marker)

al., 2018). Furthermore, an increase in *REG3G* expression with the phenotype for ETEC intestinal adhesion was seen in healthy pigs fed a low dose of oral tryptophan (17% to lysine), but not with a higher dose (22% to lysine) (Trevisi et al., 2012). In the same way, the increasing of chemokine *CCL20* expression has been linked to bacteria stimulation of pro-inflammatory signals (Skovdahl et al., 2015) while the expression of *CCL20* was reduced by the oral supplementation with antibiotic (colistin) or probiotic (*Saccharomyces cerevisiae* CNCM I-4407) (Trevisi, Latorre et al., 2017). Overall, this suggests

that apparently healthy  $MUC4^{GG}$  pigs can have a more stimulated immune system than the  $MUC4^{CC}$  pigs.

Furthermore, in our study, the *TFF3* expression was higher in the *MUC4*<sup>CG</sup> genotype than *MUC4*<sup>CC</sup> and *MUC4*<sup>GG</sup> genotypes indicating that, although *MUC4*<sup>CG</sup> and *MUC4*<sup>GG</sup> are generally considered equally susceptible to the ETEC F4ac adhesion and infection (Bijlsma & Bouw, 1987), the response in terms of gut homoeostasis of healthy pigs could be partially different between the two susceptible genotypes. Differences between the two susceptible genotypes are also

goblet cells in the mid-jejunal

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Effect of MUC4 or FUT1 polymorphisms on the score for Ulex europaeus agglutinin I (UEA) and peanut agglutinin (PNA) staining of brush border and

mucosa of pigs on d42

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TABLE

AG versus GG 0.389 0.118 0.069 0.234 0.032 0.289 AA versus GG 0.003 <0.001 <0.001 <0.001 <0.001 <0.001 AA versus AG 0.007 0.001 0.012 0.001 0.062 0.031 0.20 0.28 0.23 0.29 0.26 SEM 0.21 0.43 0.22 1.78 0.11 GG 4 က 1.25 0.70 2.53 0.57 2.96 0.71 AG Mean FUT1<sup>b</sup> 1.88 0.08 1.37 1.08 1.57 1.32 AA CG versus GG 0.878 0.862 0.703 0.617 0.695 0.467 CC versus GG 0.037 0.031 0.161 0.901 0.779 ပ္ပ CC versus 0.016 0.779 0.081 0.274 0.752 1.00 d 0.20 0.28 0.33 0.23 0.29 0.17 SEM 1.30 2.59 1.00 0.72 2.97 GG 0.8 1.15 2.65 2.78 0.92 0.72 0.85 S Ulex europaeus agglutinin I (UEA) MUC4ª Mean 0.65 1.65 2.28 0.82 0.85 S Peanut agglutinin (PNA) **Brush border Brush border** Goblet cells Goblet cells Crypts Crypts ≣> ≣

 $^{b}$ FUT1, 1-fucosyltransferase gene; genotypes had the following distribution: FUT1 $^{AA}$  = 7;FUT1 $^{AG}$  = 8; gene; genotypes had the following distribution:  $MUC4^{CC} = 8$ ;  $MUC4^{CG} = 8$ ;  $MUC4^{GG} = 8$ . as mean of assigned score values FUT1<sup>GG</sup> = 9. <sup>c</sup>Mean is expressed aMUC4, Mucin 4

supported by the results reported by Roubos-van den Hil, Litjens, Oudshoorn, Resink, and Smits (2017), which highlighted a different response in terms of ETEC F4ac faecal shedding between  $MUC4^{GG}$  and  $MUC4^{CG}$  in ETEC F4ac-infected piglets.

Furthermore, the modifications of the blood plasma metabolomics profile in pigs differing for MUC4 SNP, observed in our study. are relevant to disentangle the relationship between the genetic background of pigs and its physiological functions. Our results indicated that MUC4<sup>CG</sup> genotype had a higher level of several phosphatidylcholines with diacyl residues, which are involved in the free fatty acid metabolism, glycerophospholipid metabolism and membrane component (Vorkas et al., 2015). The MUC4<sup>CC</sup> genotype displayed a higher plasma serotonin level than the MUC4<sup>GG</sup> genotype. Serotonin is considered a neurotransmitter and paracrine signalling molecule involved in the brain-gut interactions. In our study, no signal of diarrhoea was reported; thus in this study, plasma serotonin could not be directly ascribed to the intestinal disorder. Furthermore, it is worth to mention that tryptophan is the precursor of serotonin and that a higher dietary amount of tryptophan was required for pigs with an ETEC-susceptible phenotype than not-susceptible pigs to optimize the growth response and to regulate the gene expression (Trevisi et al., 2012). Here, dietary tryptophan was provided at a sufficient level (22% to lysine), but we cannot exclude that tryptophan was limiting for plasma serotonin level in MUC4<sup>GG</sup> genotype. In the present trial, pig behaviour was not objected of specific observations; however, based on this metabolic observation and on the inverse correlation of hypothalamic serotonin with salivary cortisol (Shen, Voilqué, Kim, Odle, & Kim, 2012) and aggressive behaviour (D'Eath, Ormandy, Lawrence, Sumner, & Meddle, 2005), it will be interesting in the future to follow-up the present study considering the behaviour in growing pigs differing for MUC4 genotype.

The immunohistochemistry results for MUC4 are particularly interesting. Based on the literature, we expected an effect of MUC4 genotypes on PNA staining, since galactose has been implicated in ETEC F4ac adherence (Grange, Erickson, Anderson, & Francis, 1998; Trevisi, Miller et al., 2017), and no effect on fucosylation of the intestinal mucosa. The absence of differences in the scores for PNA in the brush borders and goblet cells of pigs for MUC4 genotype, found in our study, makes questionable the association of ETEC F4 diarrhoea with a galactose-related receptor for F4ac fimbriae. Indeed, a number of additional putative receptors for ETEC F4ac such as lactosylceramide, gangliotriaosylceramide, gangliotetraosylceramide, globotriaosylceramide, lactotetraosylceramide and lactotetraosylceramide have been described and characterized (Coddens et al., 2011; Grange, Mouricout, Levery, Francis, & Erickson, 2002). However, our data do not exclude the sialoglycoprotein (Grange et al., 1998) or n-acetyl-hexosamine nature of the receptor, where galactose may play as an enhancer of the adhesion of ETEC F4 (Grange et al., 2002). Furthermore, we cannot directly exclude the galactose implication in ETEC F4 adhesion since we analysed the intestinal PNA score of pigs under normal healthy conditions, while the galactose amount in the brush border may be affected by the ETEC presence or by

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an inflammatory bowel condition (Amador et al., 2008; Peuhkuri, Vapaatalo, & Korpela, 2010).

On the other hand, the reduced degree of fucosylation that we observed for the goblet cells in the villi in ETEC F4ac genetically susceptible pigs could be a possible indirect effect of other transfer enzymes involved in fucosylation, or maybe a consequence of a commensal bacteria stimulation able to modify the degree of mucosal fucosylation (Pickard et al., 2014).

This hypothesis is not completely supported by our results on gut microbiota composition since we did not identify any effect of MUC4 genotype for that. Our result partially contrast with the previous study by Messori et al. (2013) in which a higher presence of Clostridium bartlettii was found for MUC4<sup>GG</sup> and MUC4<sup>CG</sup> than MUC4<sup>CC</sup> pigs; however, it is generally recognized that the characterization by the sequencing of defined regions of 16S rRNA gene could have some limitations, such as an incomplete resolution and a low sensitivity (Poretsky, Rodriguez-R, Luo, Tsementzi, & Konstantinidis, 2014); thus, we cannot conclude that gut microbiota is not influenced by MUC4 genotype and further investigations are required.

On the other hand, we found significant differences in the jejunal microbiota associated with FUT1 genotypes, both in alpha and beta indices. The animals in FUT1<sup>AG</sup> group had higher internal microbial variability than the other two genotypes. Furthermore, the sPLS-DA coupled with 10-fold cross-validation revealed differences between FUT1 variants. Our results indicated that FUT1<sup>AA</sup> genotype, associated with ETEC F18 resistance, was discriminated by OTUs belonging to the Lactobacillus genus. Lactobacillus presence in the gut is mostly associated with gut eubiosis, while its reduction has been observed in case of pathogen presences and under stress condition such as weaning (Heo et al., 2013). On the other hand, the FUT1<sup>GG</sup> genotype (ETEC F18 susceptible) was discriminated by OTUs belonging to the Veillonella genus. Bacteria belonging to Veillonella genus have been recognized as non-beneficial and associated with Crohn's disease human patients (Gevers et al., 2015). Little information is reported on the FUT1 effect on intestinal microbiota. However, our results reinforce the observations reported by Poulsen et al. (2018) in which FUT1<sup>AG</sup> ETEC F18 susceptible piglets showed a higher number of non-beneficial bacteria, including haemolytic bacteria, both in faeces and in digesta, and of Enterobacteriaceae in digesta than FUT1<sup>AA</sup> piglets, even without the presences of diarrhoea. Thus, our results suggest that FUT1 genotype is not only associated with ETEC F18 resistance but can also play a role in the modulation of the commensal gut bacteria in healthy pigs. A reason of the potential relevant role of FUT1 on the microbiota could be ascribed to its functionality in the gut; FUT1 catalyses the addition of fucose to a terminal galactose in a  $\alpha$ 1,2-linkage. The mutation on FUT1 leads to a higher expression of FUT1 and FUT2 and an increase in the activity of the enzyme in FUT1<sup>GG</sup> pigs. This biological mechanism results in a different level of glycosylation on the protein structure of porcine ileum mucosa (Hesselager et al., 2016). In accordance with the immunohistochemistry observations of Coddens et al. (2007), our work showed that the FUT1AA had a lower immunoreactivity score for UEA than FUT1<sup>AG</sup> and FUT1<sup>GG</sup> and an opposite response for the PNA

immunoreactivity both in brush border and goblet cell. The *FUT1* effect on brush borders is mainly important because it represents the site where the ETEC F18 adhesion occurs. Our results contribute to explain the reason why the  $FUT1^{GG}$  genotype is more susceptible to the ETEC F18 infection. Our findings support that FUT1 susceptible pigs have a higher active fucose metabolism and display a higher level of fucose in the intestinal mucosa which can be used by the *E. coli* to increase its colonization. In addition, we observed a reverse 19 pattern for UEA and PNA immunoreactivity between the F18-susceptible or not pigs. This result can be explained by the fact that in the F18-resistant genotype, the genetically determined impaired activity of the  $\alpha(1,2)$ -fucosyltransferase could have led to the rise of the  $\alpha(1,3)$ -galactosyltransferase that can generate the typical porcine epitope  $Gal\alpha 1,3Gal$  (Phelps et al., 2003), by the glycosylation of the  $Gal\beta 1,4GlcNAc-R$  group, resulting in an increase in galactose.

The role of FUT1 in the glycosylation in the gut and on the modulation of gut microbiota could also explain the variation in the plasma metabolic profile that we observed in our study. Previous observation showed the influence of FUT1 genotypes in modulating the plasma metabolome and some specific plasma metabolites related with gut microbial metabolism (hippuric acid, oxindole, betaine) and inflammation (guanosine) (Poulsen et al., 2018). Since different techniques were applied in our and Poulsen et al. (2018) studies, and no univocal metabolites were detected, results cannot directly be compared. Here, we detected a higher amount of several phosphatidylcholines in the FUT1<sup>GG</sup> genotype than in the FUT1<sup>AA</sup> genotype. Phosphatidylcholines are indicative compounds related to inflammation and disturbed homoeostasis for their function in increasing processes of cells proliferative growth and programmed cell death (Vorkas et al., 2015). Thus, we hypothesize that gut microbiota, particularly the not beneficial genera, identified as discriminating in the FUT1<sup>GG</sup> animals, could have altered the gut eubiosis and caused stimulation of the immune system, which resulted in an alteration of the plasma metabolic profile. Furthermore, our results on gut homoeostasis in late growing phase may contribute to explain the reduced growth performance and the softer faecal score observed in the susceptible piglets during the first 3 weeks post-weaning.

In conclusion, our results improved the knowledge regarding the interplay between host genetics, gut microbiota and host physiological status. Our study indicates that MUC4 and FUT1 genotypes are not only related respectively to ETEC F4 and F18 susceptibility but can also affect the gut homoeostasis in healthy piglets in terms of fucose and galactose glycomic profiles, expression of genes related to inflammation in jejunal mucosa (MUC4) and jejunal microbiota (FUT1), and in plasma metabolic profile and growth performance. These observations could be mainly important for studies aimed to study the interplay between host and microbiota in which the genotyping of MUC4 and FUT1 genes may be used to stratify the output of studies and may contribute to a deeper understanding of host-microbial crosstalk. In addition, the early piglets' genotyping for MUC4 and FUT1 could improve the experimental protocol reliability of studies with ETEC challenge model by decreasing the number of animals needed and improving the ethicality of in vivo studies.

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#### **ACKNOWLEDGEMENT**

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#### CONFLICT OF INTEREST

The authors declare that they have no competing interests.

#### ETHICAL APPROVAL

The procedures complied with the Italian regulations pertaining to experimental animals and were approved by the Ethic-Scientific Committee for Experiments on Animals of the University of Bologna and the Italian Ministry of Health with the approval number 801/2015-PR.

#### SOFTWARE AND DATA REPOSITORY RESOURCES

The data sets on microbiota data supporting the conclusions of this article are available in the Sequence Read Archive (SRA) repository with number SRP128419.

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#### SUPPORTING INFORMATION

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