Changes in the intestinal mucosal proteome of turkeys (Meleagris gallopavo) infected with haemorrhagic enteritis virus

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A B S T R A C T

Haemorrhagic enteritis (HE) is a viral disease affecting intestinal integrity and barrier function in turkey (Meleagris gallopavo) and resulting in a significant economic loss. Sequential Windowed Acquisition of All Theoretical Fragment Ion Mass Spectra (SWATH-MS) was applied to identify crucial proteins involved in HE infection. A total of 938 proteins were identified and used to generate a reference library for SWATH-MS analysis. In total, 523 proteins were reliably quantified, and 64 proteins were found to be differentially expressed, including 49 up-regulated and 15 down-regulated proteins between healthy and HE-affected intestinal mucosa. Functional analysis suggested that these proteins were involved in the following categories of cellular pathways and metabolisms: 1) energy pathways; 2) intestine lipid and amino acid metabolism; 3) oxidative stress; 4) intestinal immune response. Major findings of this study demonstrated that natural HE infection is related to the changes in abundance of several proteins involved in cell-intrinsic immune defense against viral invasion, systemic inflammation, modulation of excessive inflammation, B and T cell development and function and antigen presentation. mRNA quantitative expression demonstrated that most of the proteins involved in innate immunity that were found to be differentially abundant were produced by intestinal mucosa, suggesting its direct involvement in immune defenses against HE infection.

1. Introduction

Haemorrhagic enteritis (HE) is an enteric disease affecting avian species, in particular turkeys (Meleagris gallopavo) older than 4 weeks of age, and compromising animal health and welfare. Haemorrhagic enteritis infection is regarded as one of the most important causes of economic loss to turkey industry. The mortality related to this disease ranges between 10–15% but it can reach 60% in some flocks (Dhama et al., 2017). The etiologic agent is the haemorrhagic enteritis virus (HE), a member of the family Adenoviridae type II (Suresh and Sharma, 1995). HE disease is characterized by acute onset of depression, pallor of the mucosa, splenomegaly, diarrhea, intestinal hemorrhage and sudden death. At day 5 after infection, haemorrhagic lesions and bleeding in the intestine start to occur. Very low levels of virus are detected in the intestinal tissue, even in the presence of intestinal lesions (Rautenschlein et al., 1998; Suresh and Sharma, 1996). Gross pathology description includes dilated intestine with blood content, and yellowish substance on the intestinal mucosa (Dhama et al., 2017). Severe congestion in the intestinal mucosa, degeneration, shortening of the villi and bleeding at the tips of them are identified at microscopic level (Sharma, 1991). Surviving birds are immunodepressed as a consequence of the lymphotrophic and lymphocytopathic attitude of the virus, which facilitates secondary infections with opportunistic pathogens agents such as the avian pathogenic Escherichia coli, bacterial respiratory diseases or septicemia (Koncicki et al., 2012; Moura-Alvare, 2014).

Although HE has been known since 1937 (Pomeroy and Fenstermacher, 1937), the molecular mechanisms of viral immunopathogenesis and immunosuppression are mostly unknown. No information about how the virus infection modifies the intestinal proteome is available as well.

To date, proteomic studies in poultry gastrointestinal tract are...
limited to chicken (*Gallus gallus*) (Luo et al., 2013; Matulova et al., 2013; O’Reilly and Eckersall, 2014; Zhang et al., 2015). To the best of our knowledge, no proteomic data about digestive protease asset is available in turkey.

In order to elucidate the molecular mechanisms underlying the pathogenesis related to HE infection, we performed a quantitative evaluation of the proteomic changes of the intestinal mucosa in turkeys after HE acute infection as compared with healthy mucosa by applying a Sequential Windowed Acquisition of All Theoretical Fragment Ion Mass Spectra (SWATH-MS) strategy. The mRNA quantification of genes related to proteins involved in immune recognition was also determined, in order to assess the effective capability of intestinal mucosa to produce the proteins that were differentially abundant.

2. Materials and methods

2.1. Samples collection and preservation

The present study was carried out on commercial B.U.T. BIG6 hybrid turkeys. Four clinically healthy turkeys were randomly selected during routine slaughtering procedures from 80 day-old females as previously described (D’Andreano et al., 2017). Pathological analysis of the gastroenteric tract evidenced no sign of gross pathological lesions related to enteritis. Molecular diagnosis to rule out the presence of HEV was carried out in the 4 tracts of intestine and spleen by means of specific PCR (Hess et al., 1999). A second group of four turkeys PCR HE positive (HE-infected animals) was included. This group of animals showed evident acute clinical signs of HE, and were subjected to euthanasia due to their critical clinical conditions, by cervical dislocation. Portions of jejunum, where the lesions were observed, were removed immediately after euthanasia: the mucosa was scratched, snap frozen into liquid nitrogen and afterwards stored at -80 °C.

2.2. Protein extraction and digestion

Protein extraction of jejunal mucosa was carried out on ice, following a general protocol for lysis of cells in 2.5 fold lysis buffer. In short, 400 mg of each tissue was washed with cold phosphate buffer saline (PBS), traces of blood were removed and tissue was homogenized in 1 ml lysis buffer (50 mM Tris pH 7.4, 150 mM NaCl, 1% Triton X-100, 1% IGEPAL, 0.1% SDS) with 10 μl Protease Inhibitor Cocktail (Sigma-Aldrich). Tissue disruption was performed using a tissue homogenizer (Precellys 24, Bertin Technologies) (4 cycles, 5500 rpm, 20 s) and glass beads (0.1 mm, Bertin Technologies) and sonicated on ice (10 cycles, 10 s). Protein extracts were centrifuged for 10 min, 10,000 x g at 4 °C. Protein concentration was determined using the method of Bradford (1976) and the absorbance was measured at 700 nm using Odyssey® CLx Infrared Imaging System (LI-COR, Biosciences).

Protein extracts were hydrolyzed in-solution, essentially as previously described (Lombart et al., 2016). Samples were adjusted to a concentration of 2 μg/μl with 6 M Urea and 50 mM NH₄HCO₃, pH = 8.5. Two μg of green fluorescence protein fused with maltose binding protein (MBP-GFP) was used as internal standard for each sample. Samples were digested with trypsin Gold (Promega) (trypsin:protein = 1:20, w/w) for 18 h at 37 °C. Each sample was completely dried and then re-suspended in 100 μl of 2% acetonitrile (ACN) with 1% formic acid (FA) and sonicated in the ultrasound bath for 5 min with low amplitude. The digested peptides were de-salted and concentrated using C18 Bond Elut OMIX solid phase extraction pipette tips (Agilent Technologies) as previously described (Silva et al., 2016). Peptides were then eluted with 100 μl of 70% ACN with 0.1% FA and, after evaporation, the pellets were stored at -20 °C.

2.3. SWATH acquisition

Peptides were resuspended in 50 μl of a solution of 2% ACN and 0.1% FA, and 10 μl of each sample were used to create two pooled samples. Four biological samples per condition were used to create the pooled samples for protein identification (four healthy animals and four HE-positive animals). In order to remove insoluble material, the peptide mixtures were then centrifuged for 5 min at 14,000 × g and collected into the proper vial for LC-MS analysis.

Samples were analysed on a Triple TOF® 5600 System (ABSciex®) in two phases: information-dependent acquisition (IDA) of the pooled samples, and, SWATH acquisition of each individual sample. Peptides were resolved by liquid chromatography (nanoLC Ultra 2D, Eksigent®) on a MicroLC column ChromXP™ C18 HL (300 μm ID × 15 cm length, 3 μm particles, 120 Å pore size, Eksigent®) at 5 μl/min with a multistep gradient: 0–2 min linear gradient from 2 to 5%, 2–45 min linear gradient from 5% to 30% and, 45–46 min to 35% of ACN in 0.1% FA and 5% dimethyl sulfoxide (DMSO). Peptides were eluted into the mass spectrometer using an electrospray ionization source (Duospray™ Source, ABSciex®) with a 50 μm internal diameter (ID) stainless steel emitter (NewObjective).

The information dependent acquisition (IDA) experiments were performed for each pooled sample in a total of two acquisitions per pool. The mass spectrometer was set to scanning full spectra (350–1250 m/z) for 250 ms, followed by up to 100 MS/MS scans (100–1500 m/z) from a dynamic accumulation time – minimum 30 ms for precursor above the intensity threshold of 1000 – in order to maintain a cycle time of 3.3 s). Candidate ions with a charge state between +2 and +5 and counts above a minimum threshold of 10 counts per second were isolated for fragmentation and one MS/MS spectrum was collected before adding those ions to the exclusion list for 25 s (mass spectrometer operated by Analyst® TF 1.7, ABSciex®). Rolling collision was used with a collision energy spread of 5.

For SWATH-MS based experiments, the mass spectrometer was operated in a looped product ion mode (Gillet et al., 2012) and the same chromatographic conditions used as in the IDA run described above. A set of 60 windows (Supplementary Table 1) of variable width (containing 1 m/s for the window overlap) was constructed covering the precursor m/z range of 350-1250. A 250 ms survey scan (m/z 350–1500) was acquired at the beginning of each cycle for instrument calibration and SWATH MS/MS spectra were collected from m/z 100–1500 for 50 ms resulting in a cycle time of 3.25 s from the precursors ranging from m/z 350 to 1250. The collision energy for each window was determined according to the calculation for a charge +2 ion centered upon the window with variable collision energy spread (CES) according to the window.

2.4. Database searches and quantification

A specific library of precursor masses and fragment ions was created by combining all files from the IDA experiments, and used for subsequent SWATH processing. The library was obtained using ProteinPilot® software (v5.1, ABSciex®), using the following parameters: i) search against a database composed by all the entries for the species *Meleagris gallopavo* (https://www.ncbi.nlm.nih.gov/protein?LinkName=genome_protein&from_uid=112) which comprised 24,245 non-redundant sequences and *Gallus gallus* (https://www.ncbi.nlm.nih.gov/protein/?term=Gallus%20gallus%20+genome) which comprised 53,468 non-redundant sequences from NCBI (April 2018), and the sequence of the recombinant MBP-GFP protein (IS); ii) iodoacetamide alkylated cysteines as fixed modification; and iii) trypsin as digestion type. An independent False Discovery Rate (FDR) analysis using the target-decoy approach provided with ProteinPilot® software was used to assess the quality of the identifications and positive identifications were considered when identified proteins and peptides reached a 5% local FDR (Sennels et al., 2009; Tang et al., 2008).

Data processing was performed using SWATH™ processing plug-in for PeakView® (v2. 0. 01, ABSciex®) (Lambert et al., 2013). Peak group confidence threshold was determined based on a FDR analysis using the
target-decoy approach and 1% extraction FDR threshold was used for all the analyses. Peptide that met the 1% FDR threshold in at least three of the four biological replicates were retained, and the peak areas of the target fragment ions of those peptides were extracted across the experiments using an extracted-ion chromatogram (XIC) window of 4 min with 100 ppm XIC width.

The levels of the proteins were estimated by summing all the transitions from all the peptides for a given protein that met the criteria described above, as adapted from (Collins et al., 2013) and normalized to the total intensity of each sample. The internal standard (MBP-GFP) was used to performed retention time alignment.

2.5. Functional annotation and grouping

The retrieved protein FASTA sequences were loaded on the open source online tool Blast2GO 4. 0. 2 software (http://www.blast2go.com). The default parameters were used and, for the basic local alignment search tool (BLAST), protein sequences were mapped against the NCBI database. The functional analysis was further narrowed by PANTHER classification (http://www.pantherdb.org). WEGO (http://wego.genomics.org.cn) has used to build graphics. Kyoto encyclopedia of genes and genomes (KEGG) database (released on April 2018) was used to classify identified proteins into specific functional terms and metabolic pathways.

2.6. RT-qPCR

In order to elucidate the capability of intestinal jejunal mucosa to produce the mRNA of proteins identified by SWATH-MS, RT-qPCR of genes involved in immunity, inflammatory pathways and response to pathogens was carried out. Total RNA was extracted from the same samples using TriZol standard protocol (Invitrogen). The RNA concentration was quantified by the NanoDrop ND-1000 spectrophotometer (Nanodrop Technologies). The purity of RNA (A260/A280) was >2. Genomic DNA was eliminated using DNase I (Invitrogen) and reverse transcription was performed with 1 μgRNA as the template, using the iSCRIPT cDNA Synthesis Kit (BioRad).

Primers were designed on turkey sequences available in NCBI by using Primer 3 (http://bioinfo.ut.ee/primer3-0.4.0), preventing possible secondary structures with the mfold Web Server (http://mfold.rna.albany.edu/?q=mfold/DNA-Folding-Form) and ensuring the specificity of the sequence by Clustal Omega (http://www.ebi.ac.uk/Tools/msa/clustalo). The primer sequences and fragment length are presented in Table 1.

Quantitative real-time PCR (qPCR) was performed using 12 μL Eva Green Supermix (BioRad) using Eco Real-Time PCR System (Illumina). The concentration of primers is reported in Table 1. GAPDH, RPL4 and YWHAZ were selected as reference genes (Marques et al., 2016). In order to assess the PCR efficiency using a relative standard curve, series of dilution were prepared by performing four-fold serial dilution starting from the pooled sample composed of a liver cDNA mix from four healthy animals. Each sample was tested in duplicate. Non-reverse transcribed controls were performed by omitting reverse transcription and no template controls were conducted by adding nuclease free water. The thermal profile used (95 °C for 10 min, 40 cycles of 95 °C for 10 s and 60 °C for 30 s; for melting curve construction, 55 °C for 15 s and 80 cycles starting to 55 °C and increasing 0.5 °C each for 10 s) was the same for each target gene. The MIQE guidelines were followed (Bustin et al., 2009). For all genes studied, the standard curves derived from serial dilution of pooled sample gave correlation coefficients (R²) greater than 0. 990 and efficiencies greater than 94%. Results were compared using the comparative Δ-ΔCq method.

2.7. Statistical analysis

Data analysis was conducted with principal component analysis (PCA) and partial least squares discriminant analysis (PLS-DA) using SIMCA P software package (Umetrics) in order to observe intrinsic clusters and obvious outliers (based on the principles of Hotelling T²) within the dataset. Statistical analysis was performed using XLSTAT for Windows (Addinsoft). Normality of data was checked by Shapiro–Wilk test. Data was compared by a Student t-test, with a two-tailed distribution and a homoscedastic variance for parametric data, while Wilcoxon signed rank test was applied in case of nonparametric data. Significance accepted for p-values ≤0.05.

3. Results

3.1. Protein identification, quantification by SWATH-MS and functional grouping of intestinal mucosa proteome

The aim of the present study was to investigate for the first time the effects of HE infection in jejunum on the proteome mapping of jejunal mucosa collected from 4 healthy and 4 HE-positive turkeys. Protein target database resources of turkey are still incomplete and poorly annotated. Therefore, aiming to increase the number of proteins that can be identified, turkey and chicken, composed by nonredundant proteins from NCBI database were combined.
Using SWATH-MS strategy, a library was created with 7013 identified peptides, corresponding to 938 different proteins across both conditions (Supplementary Tables 2 and 3, respectively). After filtering for proteins which were quantified in at least three of four biological replicates and with at least one peptide (FDR ≤ 1% and a confidence of at least 99%), this library allowed the quantification of 523 proteins among the eight samples, which were included in this comparative and descriptive analysis (Supplementary Table 4).

From a dataset composed by 523 proteins, 185 protein number identifiers were successfully functionally assigned on UniProtKB website (http://www.uniprot.org) and 45 were mapped on PANTHER website. More than 60% of the UniProtKB proteins were lacking and/or were classified as “Uncharacterized protein”; therefore, we ran BLAST using Blast2GO software to annotate the protein dataset.

Gene Ontology (GO) analysis was performed using the three main ontologies (Cellular Component, Molecular Function and Biological Process). Blast2GO retrieved 841 unique Biological Process GO terms, 541 unique Molecular Function GO terms and 800 unique Cellular Component GO terms associated with the dataset. From 523 total proteins, 18 sequences were not annotated. Gene Ontology classification using WEGO (Web Gene Ontology) Annotation Plot (http://wego.genomics.org.cn) is presented in Fig. 1.

The 523 quantified proteins were grouped into 28 GO terms divide into the three main ontology classes: i) Cellular component – 40.6% cell, 23.4% macromolecular complex, 17.2% organelle part, 10.9% extracellular region, and 7.8% membrane-enclosed lumen; ii) Molecular function – 57.8% binding, 39.1% catalytic activity, 3.1% transporter activity; and iii) Biological process – 42.2% metabolic process, 42.2% cellular process, 6.3% localization, 4.7% pigmentation, 3.1% anatomical structure formation, and 1.6% response to stimulus.

3.2. Differential quantification and statistical analysis of intestinal mucosa proteins between HE and control groups

To identify differentially abundant proteins between the two studied conditions, statistical analysis was applied, and fold-changes and p-values were used to rank and filter the quantitative data (Fig. 2).

Differentially abundant proteins were defined as those that showed a fold change with a p-value ≤ 0.05. In total, 64 proteins differentially abundant between the healthy and HE groups were identified in the turkey intestine, which included 49 proteins up-regulated and 15 down-regulated proteins in HE-affected animals (Table 2). Following KEGG classification, the 64 differentially abundant proteins were grouped into nine classes based on their putative functions: amino acid and protein metabolism (25%), carbohydrate metabolism (19%), fatty acids metabolism (16%), cytoskeleton (11%), immune system (11%), nucleic acid biogenesis (6%), ribosome (6%), binding and transport (5%) and apoptosis (1%), as shown in Fig. 3. Proteins related to amino acid and protein metabolism, carbohydrate metabolism, fatty acids metabolism, cytoskeleton and immune system were predominant and accounted for approximately 80% of the differential proteins.

The significant similarity of the 4 biological replicates samples and the noticeable differences of protein patterns between healthy and HE-infected intestine did not result in distinct segregation of the proteomes after PCA analysis applied to the 523 quantified proteins as multivariate statistical method. On the contrary, four healthy animals and four diseased animals show a clear separation into two groups after PLS-DA analysis (Fig. 4).

3.3. mRNA expression of selected proteins related with immune system

The intestine is crossed by a wide capillary network. The finding of a protein by proteomic techniques does not confirm its expression by intestine. Therefore, several proteins can be expressed by liver or other tissues and then delivered to intestine through blood. The effective ability of jejunal mucosa to produce proteins related with immune response was confirmed by RT-qPCR through detection of their respective mRNAs. We focused our attention on those differentially expressed proteins that were found to be up-regulated more than 2 times in healthy mucosa and involved in innate immunity and inflammatory response, namely selenium-binding protein 1-A (SELENBP1), heme-pexin (HPX), PIT54, fibrinogen beta chain (FGB), membrane-associated progesterone receptor component (PGRMC1), immunoglobulin lambda-like polypeptide 1 (IGLL1) and cell surface A33 antigen (GPA33),

![Fig. 1. Proteome characterization of mucosa intestine. GO classification using Blast2GO shows level 2 categories for Cellular Components, Molecular Function and Biological Processes.](image-url)
coronin-2A (CORO2A), 3-mercaptopypyruvate sulfurtransferase (MPST), calnexin (CANX), peptidyl-prolyl cis-trans isomerase A (PPIA) and calmodulin (CALM), which are related to inflammation, viral and bacterial defenses and response to external stimulus.

The mRNA expression studies are presented in Fig. 5. Quantitative analysis confirmed that at least 6 of the proteins involved in immunity and inflammatory response were produced by jejunal mucosa, although the difference between healthy and HE infected animals was found not significant. mRNA coding for SELENBP1, PPIA, MPST, PGRMC1, CANX and CALM didn’t show any statistical difference. Remarkably, the relative differences in abundance between amount of the three acute phase proteins (APP) included in the list of differentially expressed proteins, namely FGB, PIT54 and HPX, is negligible in both healthy and infected mucosa.

4. Discussion

This study shows the differential changes in the intestinal proteome of turkey and how it is modified following the infection with HE. Although HE is regarded as one of the leading causes of viral intestinal diseases in turkeys, the molecular background of its pathogenesis and how the local immune system reacts to the invasion of the virus are mostly unknown. By using a next generation proteomic approach, the SWATH-MS, we described the jejunal mucosa proteomes of HE-infected turkeys as compared to healthy ones. The effective capability of jejunal mucosa to produce the proteins identified by means of proteomics analysis was further investigated by quantitative gene expression analysis.

The KEGG classification analysis revealed that out of 64 differentially abundant proteins in HE intestinal mucosa proteome, a number of 43 (66%) are involved in protein, carbohydrate, fatty acid and nucleic acid metabolism. This result is consistent with previous metabolic studies showing that viruses can induce modification in host cell metabolism, including an increase of glycolysis and fatty acid synthesis, increasing the amount of energy available for virus replication (Thai et al., 2014). Changes in nucleotide, amino acid and protein metabolism may provide an increase in pools of free aminoacids and nucleotides necessary to cope with viral genome replication and virus assembly (Sanchez and Lagunoff, 2015). A total of 11% of differentially abundant proteins were related to cytoskeleton, including, among the others, WD and Coronin1, which are decreased in intestinal mucosa from HEV affected turkeys. Coronins are essential regulators of Ca²⁺ trafficking after T cell receptor stimulation, and a decrease of Coronin1 has been linked to a CD4⁺ T cell response during viral infection (Tchang et al., 2013).

The focus of the present work was to study the involvement of intestinal mucosa on immune defenses and inflammation against HE infection. We provided the evidence that natural HE infection is linked to the changes in abundance of proteins related to cell-intrinsic immune defense against viral invasion, systemic inflammation, modulation of excessive inflammation, B and T cell development and function and antigen presentation.

4.1. Proteins related to cell intrinsic immune defense against virus

HE infection induces a decrease in the abundance of DDX3X (DEAD-Box Helicase 3 X-Linked). Decreasing of DDX3X leads to a significant reduction of infectious HSV-1 particles (Khadijvjam et al., 2017), suggesting the activation of a defensive mechanism in HE as well.

4.2. Proteins involved in systemic reaction to inflammation

As expected, HE infection prompted an inflammatory response as confirmed by the increased abundance of proteins related to systemic reaction to inflammation, such as FGB (fibrinogen β chain), PIT54 and HPX (Hemopexin) (O’Reilly and Eckersall, 2014; Adler et al., 2001; Ceciliani et al., 2002). The presence of FGB in turkey HE-intestine confirms previous studies in chicken infected with gastrointestinal diseases (Georgieva et al., 2010). The PIT54 is regarded as one of the major APP in poultry (O’Reilly and Eckersall, 2014) and is upregulated in turkeys stressed after road transport (Marques et al., 2016). PIT54 expression has been recently demonstrated in chicken, but its mRNA abundance in healthy animals was shown to be negligible (Marques et al., 2017). The increased abundance of FGB, PIT54 and HPX provides the evidence that HE infection triggers a local inflammatory reaction and, given the involvement of fibrinogen, the activation of a coagulation cascade. On the background that the increase of mRNA of FBG, PIT54 and HPX in intestinal mucosa of HE infected turkeys is negligible, we hypothesize that the local increase of APP is related to a translocation of proteins from blood serum to intestinal mucosa. These hypothesis is confirmed by previous finding (Takagi et al., 2012) that have also reported the presence of HPX protein in

Fig. 2. The distribution of p-values and fold changes (log10) in 523 quantitative proteins between the healthy and HE-infected groups. A total of 64 proteins were selected as different proteins, which exhibited a p-value ≤ 0.05.
injured intestinal mucosa of rats, but not at mRNA level.

### 4.3. Proteins involved in changes of B-cell and T-cell population

One of the major features of HE infection is turkey is a dramatic change of B-cell and T-cell population, which is depleted because of the viral infection ([Suresh and Sharma, 1996, Suresh and Sharma, 1995]). The proteomic results presented in this study partially provide a molecular background to these changes, demonstrating that HE infection is related to changes in the abundance of proteins involved in B-cells and T cells development and differentiation, such as IGLL1, PGRMC1, PPIA and CALM.
The IGLL1 gene encodes a component of the pre-B-cell receptor, which is crucial for B cells development (Chen et al., 2016) and dysfunctions of the IGLL1 gene cause a primary immunodeficiency related to poorer proliferation and differentiation of pro-B cells and consequently, lower levels of serum antibodies and circulating B cells (Bankovich et al., 2007). PGRMC1 is a single transmembrane protein that forms part of a multi-protein complex that binds to progesterone and other steroids, and a recent report suggested that its expression is

Fig. 3. Functional classification of the proteins of differential abundance identified from the intestinal mucosa of turkey.

Fig. 4. PCA and PLS-DA score-plots from healthy and HE-affected groups. A: Score-plot obtained from PCA modelling, 2 components extracted. Healthy group, green. HE-infected group, blue. The first component explains 35% ($R^2X[1] = 0.345$) of the variation and the second component 17% ($R^2X[2] = 0.167$). B: Score-plot obtained from PLS-DA modelling, 2 components extracted. Healthy group, green. HE-infected group, blue. The first component explains 30% ($R^2X[1] = 0.269$) of the variation and the second component 20% ($R^2X[2] = 0.20$). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article).
involved in Th1/Th2 and Treg regulation (Maeda et al., 2013). PPIA (also known as Cyclophilin A) is associated with viral infections in animals. Folds relative to liver = 1. Relative expression of PIT54, PGRMC1, SELENBP1, HPX, FGB, CANX, MPST, PPIA and CALM in liver and intestine of healthy and HE-infected turkeys were studied by qPCR. The results were normalized using the geometric mean of MPST, PPIA and CALM in liver and intestine of healthy and HE-infected turkeys.

4.4. Proteins involved in modulating excessive immune response

The proteomic analysis carried out in this study also identified a significant increase of proteins whose role is to modulate excessive immune response, such as SELENBP1, COROA2 and MPST. SELENBP1 is a member of selenium-binding protein family, which has been shown to bind selenium covalently (Porat et al., 2000). The role of SELENBP1 in intestine is to modulate the differentiation and function of immune cells, contributing to reduce excessive immune response (Speckmann and Steinbrenner, 2014). COROA2 has been shown to fulfill an anti-inflammatory function, being required to de-repress TLR target genes by macrophages (Huang et al., 2011). Further protective effects against excessive inflammatory response may be provided by MPST (Mercaptopyruvate Sulfurtransferase), whose abundance is increase in HE affected animals and may provide an antioxidant activity (Nagahara et al., 2019).

4.5. Proteins involved in antigen presentation

The list of proteins whose abundance is modified due to HE infection in intestinal mucosa includes also proteins involved in antigen presentation, such as Calnexin (CANX) and GPA33. CANX is part of the chaperon machinery involved in antigen presentation and is also involved in intestinal Paneth cells’ differentiation (Gassler et al., 2002; Huang et al., 2016). GPA33 is also involved in immune signaling and antigen presentation. GPA33 is an intestinal epithelium-specific cell surface marker and a component of the tight junction-associated proteins of the immunoglobulin superfamily (Ackerman et al., 2008). Beside its activity on antigen presentation, GPA33 is also crucial for the maintenance of intestinal barrier function (Williams et al., 2015). An increased abundance of GPA33 might be related to a protective function against excessive intestinal leakage.

5. Conclusions

This study presented the first proteomic comparison of healthy and HE-affecte intestinal mucosa proteome in turkey, focusing on immune reaction against virus, and demonstrating that several proteins that were found to be differentially expressed are involved in several pathways related to immune defense against viruses. Our findings provide insight in the immunological pathways activated by HE infection, in particular for what concerns the different abundance of proteins involved in B and T cell regulation and development. Further work is necessary to establish the precise mechanism by which up-regulating or down-regulating of these proteins provide protection against the disease.

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Appendix A. Supplementary data

Supplementary material related to this article can be found, in the online version, at doi:https://doi.org/10.1016/j.vetimm.2019.06.001.

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