High dosage of zinc modulates T-cells in a time-dependent manner within porcine gut-associated lymphatic tissue

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(Submitted 14 February 2018 – Final revision received 1 August 2018 – Accepted 12 September 2018)

Abstract

Zn serves as a powerful feed additive to reduce post-weaning diarrhoea in pigs. However, the mechanisms responsible for Zn-associated effects on the adaptive immune responses following feeding of a very high dosage of Zn remain elusive. In this study, we examined the T-cell response in gut-associated lymphatic tissues of seventy-two weaned piglets. Piglets received diets with 57 mg Zn/kg (low Zn concentration, LZn) or 2425 mg Zn/kg (high Zn concentration, HZn) mg Zn/kg feed for 1, 2 or 4 weeks. We observed that feeding the HZn diet for 1 week increased the level of activated T-helper cells (CD4+ and CD8+T-cells) compared with the MZn and LZn groups (P < 0.05). In addition, we observed higher transcript amounts of interferon γ and T-box 21 (TBET) in the HZn group compared with the MZn and LZn groups (P < 0.05). A gene set enrichment analysis revealed an over-representation of genes associated with cytokine signalling in immune system. Remarkably, feeding of a very high Zn dosage led to a switch in the immune response after 2 weeks. We detected higher relative cell counts of CD4+CD25(bright) regulatory T-helper cells (CD4+ T-cells) and a higher expression of forkhead box P3 (FOXP3) transcripts (P < 0.05). After 4 weeks of feeding a high-dosage Zn diet, the relative CD4+ T-cell count (P < 0.05) and the relative CD8β T-cell count (P < 0.01) were reduced compared with the MZn group. We hypothesise that after 1 week the cellular T-helper 1 response is switched on and after 2 weeks it is switched off, leading to decreased numbers of T-cells.

Key words: Zinc: Nutritional immune response: T-cells: Pigs

The micronutrient Zn is an essential trace element that is required for physical growth, development and immune defence(1). A lack of Zn can lead to growth and developmental retardation, an impairment of the immune system together with decreased resistance to infections and therefore can cause diverse diseases in humans and animals(2–5). Several disorders related to Zn deficiency such as diarrhoea have been shown to be recovered and prevented by Zn supplementation as adjunct therapy(6). Different studies demonstrate that Zn deficiency impairs the formation and activity of inflammatory cytokines and therefore has an influence on the development and regulation of immune cells(4,5). A deficit of Zn can thus alter the immune response, resulting in an ineffective control of pathogens(6).

However, further investigations are needed to fully understand the precise effects of Zn on the immune system. In particular, the pig, which is known to have a digestive physiology similar to humans as it is an omnivore and the organ size is comparable to humans, could be used as a translational model(7,8).

Abbreviations: BW, body weight; FOXP3, forkhead box P3; GALT, gut-associated lymphatic tissue; GATA3, GATA binding protein 3; GSEA, gene set enrichment analysis; HZn, high Zn concentration; IFNγ, interferon γ; LZn, low Zn concentration; MLN, mesenteric lymph node; MZn, medium Zn concentration; TBET, T-box 21; ZIP4, solute carrier family 39 member 4 (SLC39A4); ZnT, solute carrier family 30 (SLC30).

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In pig husbandry, Zn is known to induce several positive effects on the gastrointestinal microbiota when fed at a high dietary level (2000–3000 parts per million (ppm)), including enhanced growth of the pigs, and shows improved resistance against post-weaning diarrhoea\(^{(9,12)}\).

Despite its positive effects, there are also negative aspects of Zn that need to be considered. Several lines of evidence suggest that long-term uptake of high dosage of Zn can lead to negative effects on the immune system. Previous studies using an infection model with *Salmonella typhimurium* revealed that long-term application of zinc oxide over a period of 6 weeks led to decreased number of T-cell populations within the mesenteric lymph nodes (MLN) of the pigs\(^{(13)}\). Within the same experimental setup, we observed increased expression of suppressor of cytokine signalling (SOCS) SOCS7 in MLN of piglets, which suggested that such treatments may impair the immune response\(^{(14)}\). These results are consistent with previous findings that permanently high uptake of Zn can result in an impairment of lymphocyte proliferation\(^{(15)}\). Moreover, there is some evidence for elevated levels of liver enzymes associated with liver damage in piglets\(^{(16)}\) supplemented with high level of Zn, and co-accumulation of Cu and Zn was found in the kidney of piglets fed very high concentrations of zinc oxide for 4 weeks\(^{(17)}\). It was shown that the CD8\(^{+}\) γδ T-cell fraction of intra-epithelial lymphocytes showed reduced abundance after 4 weeks of feeding a high Zn diet compared with the control diet\(^{(18)}\). However, a more detailed analysis of the immune-regulating genes providing insight into beneficial and impairing mechanisms of the T-cell immune response within the gut-associated lymphatic tissues (GALT) after feeding high Zn dosages is still missing.

**Sampling of gut-associated lymphatic tissue and immune cells**

In all, eight piglets per group and time point were euthanised at the following time points: 32 (SD 1), 39 (SD 1) and 53 (SD 1) d of age. The piglets were sedated with 20 mg/kg BW of ketamine hydrochloride (Ursotamin\(^{®}\), Serumwerk Bernburg AG) and 2 mg/kg BW of azaperone (Stresnil\(^{®}\), Jansen-Cilag), and cord blood was taken before euthanasia with intra-cardial injection of 10 mg/kg BW of tetracaine hydrochloride, mebezonium iodide and embutramide (T61\(^{®}\); Intervet). During necropsy, jejunal and ileocaecal mesenteric lymph nodes (JELN and ILLN, respectively) and Peyer patches from the jejunum and the ileum (JEPP and ILPP, respectively) were collected and either used for RNA isolation or weighed. In the latter case, tissue weights were expressed as milligrams per 100 g of tissue. All lymphoid tissues were freeze-dried before hydrolysis in hydrochloric acid (12 M) and subsequently analysed by atomic absorption spectrometry in an AAS vario 6 spectrometer (Analytik Jena). We aimed to have 50, 150 or 2500 mg Zn in 1 kg of feed; the actual measured values for the feedings groups were 57 (LZn), 164 (MZn) or 2425 (HZn) mg Zn/kg. The Zn source was analytical-grade zinc oxide (Sigma Adrich). No antibiotics were administered before and during the experiment. All animal experiments were performed at the Institute of Animal Nutrition, Freie Universität, Berlin, Germany, in accordance with the German Animal Protection Law and approved by the local responsible authorities (LaGeSo Reg. No. 0347/09). The *in vivo* procedures were consistent and in compliance with the Animal Research: Reporting of *In Vivo* Experiments (ARRIVE) Guidelines for Reporting Animal Research.

**Methods**

**Animals, housing and diets**

A total of seventy-two purebred Landrace piglets were weaned at 26 (SD 1) d of age with a mean body weight (BW) of 7.2 (SD 1-2) kg and randomly allocated into three dietary treatment groups balancing for sex, litter and BW. BW and lymphoid tissue sizes can be found in online Supplementary Table S1; pairwise *t* tests were performed between the three feeding groups to detect difference in relative lymphoid tissue size between the groups (no significant differences were detected; online Supplementary Table S1 – sheet 4). Animals were housed in pens (2 per pen) with straw bedding and *ad libitum* access to feed and water. Each dietary group (*n* 24 per feeding group) was fed a common basal maize–wheat–barley–soyabean diet with different Zn levels (online Supplementary Table S2)\(^{(23)}\). The dietary Zn levels were confirmed by analysis using atomic absorption spectrometry in an AAS vario 6 spectrometer (Analytik Jena). We aimed to have 50, 150 or 2500 mg Zn in 1 kg of feed; the actual measured values for the feedings groups were 57 (LZn), 164 (MZn) or 2425 (HZn) mg Zn/kg. The Zn source was analytical-grade zinc oxide (Sigma Adrich). No antibiotics were administered before and during the experiment. All animal experiments were performed at the Institute of Animal Nutrition, Freie Universität, Berlin, Germany, in accordance with the German Animal Protection Law and approved by the local responsible authorities (LaGeSo Reg. No. 0347/09). The *in vivo* procedures were consistent and in compliance with the Animal Research: Reporting of *In Vivo* Experiments (ARRIVE) Guidelines for Reporting Animal Research.

**Zinc concentration in mesenteric lymph nodes**

MLN were freeze-dried before hydrolysis in hydrochloric acid and subsequently analysed by atomic absorption spectrometry in an AAS vario 6 spectrometer. An external standard (PerkinElmer Pure XVI, #N9300281; PerkinElmer) was used for calibration. The accuracy was checked by a certified reference Zn solution (Titrisol, #9953; Merck).
Cell culture of isolated leucocytes from mesenteric lymph nodes

As direct effects of Zn on the immune cells are difficult to study in vivo, owing to many environmental influences of immune status, we opted for a cell culture approach using primary isolated immune cells. We believe that this approach should lead to reduced complexity of the studied system compared with in vivo experiments. However, the use of primary isolated immune cells should give results reflecting the interplay of different immune cells more accurately compared with using single-origin cell lines. Cells were isolated from one healthy control piglet, 1 × 10^6 cells/well were cultured in a ninety-six-well plate using Roswell Park Memorial Institute (RPMI) media enriched with 10% fetal calf serum, 200mM l-glutamine and Antibiotic–Antimycotic (Thermo Fisher Scientific). The aim of the in vitro experiments was to mimic the same concentrations of Zn in cell culture as they occur in vivo. The analyses of Zn concentrations in MLN revealed on average 13 mg Zn/kg. If one accounts for the molarity of Zn, this calculates 0.2 mM Zn/litre in the cell medium to reflect the in vivo measured concentration. However, this is not the concentration in the cell. In pre-experiments, we tested Zn concentrations from 10 μM to 2 mM from different organic and inorganic Zn sources in porcine and murine splenocytes and lymphocytes from MLN. Although ZnSO₄ and zinc oxide were toxic to the primary cultured cells in the used concentrations of 100 or 200 μM Zn²⁺, the cells survived for 5 d when the medium was supplemented with 200 μM zinc gluconate or zinc acetate per litre. For our experiments, we chose to add zinc acetate and we calculated the Zn²⁺ amount to be 10, 100 or 200 μM Zn²⁺ per well to reflect low and high levels of Zn for the primary cultured leucocytes.

Flow cytometry

Flow cytometry (FCM) was performed with the purified lymphocytes from the intestinal tissues and blood. Combinations of surface antigens were used to detect the following cell types – T-helper cells: CD4⁺ (Clone 74-12-4; Southern Biotech), CD25⁻/⁻ (Clone K231.3B2; Biozol) and CD8α⁻/δ-∆in (Clone 76-2-11; Southern Biotech); regulatory T-cells: CD4⁺, CD25⁺high, and cytotoxic T-cells: CD8β¹ (Clone: PG164A; VMRD) and CD4⁻. For forty randomly chosen samples, we performed an additional intra-nuclear staining of forkhead box P3 (FOXp3) (Clone FJK-16s; eBioscience) to assess the regulatory T-cell phenotype as well. In these samples, the relative cell counts of CD4⁺CD25⁺high and CD4⁺FOXp3 were the same. For each reaction, 1 × 10⁶ cells were stained as described elsewhere. Per sample, 50,000 lymphocytes, which were negative for propidium iodide staining (0.5μg/ml), were assessed by FCM using a BD FACSCalibur™ flow cytometer (Becton Dickinson).

Gene expression in tissue and isolated lymphocytes

Tissue samples were homogenised by the FastPrep-24 Instrument (MP Biomedicals), and RNA was isolated using the NucleoSpin® RNA II Kit of Macherey-Nagel according to the manufacturer’s instructions. The integrity of total RNA was measured by the Agilent 2100 Bioanalyzer (Agilent Technologies). RNA samples with an RNA integrity number (RIN) value higher than 7.5 were taken for quantification. The quantity was assessed by a NanoDrop spectrophotometer (PEQLAB Biotechnologie GmbH). In all, 1 μg of total RNA was reverse-transcribed into complementary DNA (cDNA) using AffinityScript quantitative PCR (qPCR) cDNA Synthesis Kit (Agilent) with oligo(dT) primers. Expression of 60S ribosomal protein L19 (RPL19) and TATA box-binding protein (TBP) did not vary owing to the Zn treatment of the piglets within the used tissues and cells and were therefore used as stable reference genes for normalisation of expression data for every sample on each plate. The primers and their source are described in online Supplementary Table S3. Primers for T-box 21 (TBET), FOXP3, GATA binding protein 3 (GATA3), interferon γ (IFNγ), IL12 and protein kinase C theta (PKCθ) as well as the solute carrier family 39 member 4 (SLC39A4; ZIP4) transcripts were designed with exon-boundaries overlapping, based on published sequences of the pig (Ensembl, Genome assembly: Sscrofa10.2) using the online primer design tool Primer3 (http://frodo.wi.mit.edu). Expression of the Zn transporter genes ZnT1, ZnT2 and ZnT5 (solute carrier family 30; SLC30) was determined for the principal isoform. The Zn transporter gene ZIP4 was examined for different protein-coding transcripts. The long transcripts ZIP4-201 and ZIP4-001 contain all twelve exons of the ZIP4 gene; however, ZIP4-001 has a shortened exon 1 with a back-shifted transcription start site. The real-time qPCR (RT-qPCR) reaction was carried in a total volume of 10 μl containing 6 μmol forward primer, 6 μmol reverse primer, 10 ng of cDNA and 5 μl of SYBR Select Master Mix (Applied Biosystems). The amplification protocol included a 10-min denaturation step at 95°C, followed by forty cycles consisting of 30 s at 95°C, 20 s at 60°C and 40 s at 72°C. The quantification of the transcript amounts was performed with a ViiA™ 7 Real-time PCR System (Applied Biosystems) or for the cell culture experiments with a Thermal Cycler (Bio-Rad). Afterwards, a melting curve analysis was performed in every reaction well to check for specificity of primers. All primer sets were initially validated for single amplicon generation, and standard curves were tested before analysis. Only primer pairs with efficiency rates between 90 and 110% were considered for gene expression analyses. Melting curves and PCR efficiency were used as standard quality criteria for each RT-qPCR run. Relative transcript amounts were calculated using the relative quantification method (ΔΔCt). To calculate the ΔCt, the mean threshold cycle (Ct) values of the two endogenous control genes RPL19 and TBP were subtracted from the Ct of the target gene. To compare the relative expression among the different tissues and diets, the mean of the ΔCt values of animals belonging to the MZn group (our control group) from the Peyer’s patch were taken as the external calibrator to calculate the ΔΔCt. Results are presented as relative expression = 2⁻ΔΔCt. Results shown for GALT represent ΔΔCt values of animals belonging to the MZn group and not the external calibrator. The ΔΔCt values of the piglets from the Peyer’s patch were chosen to be the external calibrator to calculate the ΔΔCt. Results are presented as relative expression = 2⁻ΔΔCt. Results shown for GALT represent ΔΔCt values of animals belonging to the MZn group and not the external calibrator. The ΔΔCt values of the piglets from the Peyer’s patch were chosen to be the external calibrator.
approach for mRNA of MLN from 32-d-old piglets after feeding MZn or HZn in the feed for 1 week. Sequencing of individually barcoded samples was carried out for two male and one female piglet for MZn and HZn.

Read alignment. Paired-end sequencing reads were obtained using an Illumina instrument and were trimmed using trimomatic 0-32 using the TruSeq3-PE-2.fa adapter library from Illumina. After trimming alignment was done by tophat2 using the Sscrofa10.2 genome (Ensembl) and the Sscrofa10.2 transcriptome (Ensembl), we did not allow for multi-mapped reads (prefilter-multihits) and allowed for two mismatches, 13-bp alignment gaps and a maximum insertion and deletion size of 20bp when aligning sequences to the reference genome/transcriptome. After alignment reads were sorted using samtools (version 1.3.1-45) and optical duplicate were marked using picard tools (version 1.99), the resulting bamfiles were indexed using samtools, after which base recalibration around known SNP was performed using GATK (3.2-2) according to GATK best practices.

Gene expressions from RNA-sequencing. After alignment of reads, read-level reads per kilobase million (RPKM) gene expressions were performed using the following procedure: using the GenomicFeatures R package, we made a transcript database (exons by gene) from the Sus_scrofa.Sscrofa10.2.84. short.gtf file obtained from Ensembl. Read overlap between the genes and bamfiles was performed using the summarizeOverlaps function from the GenomeAlignments R package in union mode, ignoring strand orientation but allowing for fragments within a feature to be counted. Raw read counts were extracted and quantile normalised and transformed into RPKM values. These data were then log2-transformed to follow a normal distribution followed by differential expression analysis using standard two-sided t tests. Genes were annotated using biomaRt, using the sscrofa_gene_ensembl database (online Supplementary Table S4).

Gene set enrichment analysis. We used the FGSEA package in R to perform gene set enrichment analysis (GSEA) on our differentially expressed genes. Therefore, we converted the Ensembl gene identifiers of our data into Entrez identifiers using the biomaRt R package. When a gene had no corresponding Entrez identifier in the biomaRt conversion, we used the Entrez identifier associated with the human orthologue for the corresponding gene. After converting identifiers, we then used the reactomePathways function provided by the FGSEA package to map pathways annotations to our Entrez identifiers. Settings used in the fgsa analysis function were 10 000 000 permutations; minimum size of the gene set to test was 80; and the maximum size of the gene set to test was 500. These parameters were chosen as we only want to look for over-representation in larger pathways as our sample size is low, and we expect a lot of false positives when performing GSEA using small pathways. Finally, we extracted over-represented pathways and plotted the figure using the topPathways function.

Statistical analyses

Statistical analyses of data. FCM raw data were analysed using FlowJo version 7.9 and further processed using R version 3.2.3. Plots were generated using the R package ggplot2. Zn group pairs (LZn against MZn and HZn against MZn) were tested using pairwise Mann–Whitney U tests. Multiple testing correction was not applied because of the small sample sizes. Relative gene expressions were calculated and analysed with Microsoft Excel 2010. Results were considered significant when P ≤ 0.05. The effect of Zn concentrations was tested using ANOVA (Table 1); as no significant differences were detected, a post hoc test was not performed.

Statistical power estimation. Estimation of differences in relative cell count, as well as in relative expression of genes, using a statistical power of 80% is performed using pairwise Whitney U tests for the LZn and HZn dosage groups against the control MZn group. The minimally detectable effect size required was set to 1 unit of difference in relative expression level or relative cell count, leading to an estimated eight samples required per group. When three time points were considered (1, 2 and 4 weeks), we estimated in total to detect significant effects with seventy-two pigs.

For RNA sequencing analysis, six animals were used (three HZn v. three MZn), and this was done owing to monetary limitations, as we do not expect to reach a power of 80% at a significance level of 0.05 for testing differentially expressed genes directly from these data. Therefore, we used the data from RNA sequencing to perform GSEA. GSEA can be used to pick up general tendencies in the expression data and point to affected gene sets/pathways that are effected by Zn concentration (HZn v. MZn). Unfortunately, it is not possible to estimate the power obtained by GSEA; however, we control results at a false discovery rate of 5%.

Results

Expression of zinc transporters and zinc concentration in gut-associated lymphatic tissue

The expression of Zn transporters is a prerequisite for the proper uptake of Zn ions by immune cells and immune system-associated cells. We explored the repertoire of Zn transporters of GALT of piglets fed LZn, MZn and HZn diets for 1 and 2 weeks (Table 1). The expression of the long ZIP4 transcripts ZIP4-201 and ZIP4-001 and the Zn transporters ZnT1, ZnT2 and ZnT5 was measured in the MLN and Peyer’s patches. The data provide evidence that ZIP4, ZnT1, ZnT2 and ZnT5 are expressed not only in the epithelium but also in the MLN and Peyer’s patches of the jejunum (Table 1). However, there was no significant expression difference in these lymphoid tissues among the feeding groups for any examined Zn transporter gene, although we found a slight increase in Zn (14-31 mg/kg) in the HZn group compared with MZn (11-55 mg/kg; P < 0.05) and LZn (12-45 mg/kg; P < 0.1) groups in MLN after 1 week of feeding the Zn diets (Fig. 1). After 4 weeks of feeding, there was no difference in the Zn concentration for the feeding groups.
Table 1. Influence of dietary zinc concentrations low zinc concentration (LZn), medium zinc concentration (MZn) and high zinc concentration (HZn) on the relative expression of the zinc transporter transcripts in Peyer's patches and lymph nodes of the jejunal small intestine of piglets after 2 weeks of feeding* (Mean values and standard deviations)

<table>
<thead>
<tr>
<th>Gene</th>
<th>Tissue</th>
<th>Mean</th>
<th>SD</th>
<th>n</th>
<th>LZN</th>
<th>Mean</th>
<th>SD</th>
<th>n</th>
<th>MZN</th>
<th>Mean</th>
<th>SD</th>
<th>n</th>
<th>HZN</th>
<th>Mean</th>
<th>SD</th>
<th>n</th>
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<tr>
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<td>3.01</td>
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<td>±1.56</td>
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<td>±1.35</td>
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ZIP4, solute carrier family 39 member 4 (SLC39A4); ZnT, solute carrier family 30 (SLC30).

* LZn (low dietary Zn) = 57 mg Zn/kg feed; MZn (medium dietary Zn) = 164 mg Zn/kg feed; HZn (high dietary Zn) = 2425 mg Zn/kg feed. All expression values of the transcripts are relative to the mean of the ΔCt values of the MZn group from the Peyer’s patch.

Short-term effects of zinc on T-cells

After 1 week of feeding a HZn diet, piglets had a higher relative cell count of CD4+ and CD8α dim T-cells than piglets fed diet with a low concentration of Zn (LZn) (P < 0.05). Hence, we could confirm the in vitro results in vivo. Within the cell culture-based experiment, we found a tendency of lower transcript amounts of FABP5 in the 100 and 200µM Zn2+ treatment groups compared with the control and the LZn treatment groups. After 1 week of feeding the different Zn diets, the T-helper cell-associated genes TBET, GATA3 and FOXP3 were up-regulated at the transcript level in the HZn group when compared with the LZn (P < 0.05) and MZn (P < 0.05) groups within the GALT (Fig. 2(c)). Consequently, we performed in vitro assays, using isolated immune cells. In these in vitro assays, using isolated lymphocytes from the cell culture-based experiment. After 3 d of incubation with 0, 10, 100 or 200µM Zn2+, we observed a trend with 10µM Zn2+ when compared with the untreated controls, after 1d of treatment of CD4+ and CD8α dim T-cells with 100 and 200µM Zn2+. After 4 weeks of feeding a HZn diet, piglets had a tendency of lower transcript amounts of TBET, GATA3 and FOXP3, also in the jejunal lymph nodes compared with the control and the EZn (P < 0.05) and MZn (P < 0.05) groups within the GALT (Fig. 2(c)). Consequently, we performed in vitro assays, using isolated immune cells. In these in vitro assays, using isolated lymphocytes from a healthy donor, piglets which were treated with 1μM Zn2+ for 3 d, we detected a trend with 100 and 200µM Zn2+ , we observed a higher transcript amounts of TBET, the master transcription factor of T-helper cells, FOXP3 and GATA3 (P < 0.05) when compared with the untreated controls, after 1d of treatment of CD4+ and CD8α dim T-cells with 100 and 200µM Zn2+. 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treatment group of 10 µM Zn²⁺ (Fig. 2(d)). GATA3 and FOXP3 showed no difference in expression after short-term feeding of zinc oxide (data not shown). In addition, we set up an in vitro experiment performed in a different laboratory using a different Zn source, zinc gluconate, to validate the found Zn effects on TBET within MLN. We also detected a higher expression of TBET 100 and 200 µM Zn²⁺ treatment groups compared with the control and the Zn treatment group of 10 µM Zn²⁺ (online Supplementary Fig. S1). Furthermore, we checked the response of lymphocytes derived from blood within the in vitro assay using zinc gluconate. We could not detect a difference in TBET expression for lymphocytes from blood. In a next step, we were interested in the expression of important T-regulating genes and choose IL12 and PKCβII. Remarkably, PKCβII was highly up-regulated with Zn treatment in MLN-derived cells, whereas the up-regulation in blood-derived immune cells was moderate. IL12 showed an increase of expression in lymphocytes derived from MLN and from blood in the HZn treatment groups.

To better understand the mechanism, accounting for the short-term effects of high dietary Zn, we performed RNA sequencing of MLN from 32-d-old piglets after feeding MZn (MZn= three individual piglets) and HZn concentration (HZn= three individual piglets) in the feed for 1 week. To get a picture of contributing pathways, we performed GSEA on our differentially expressed genes. GSEA revealed highest enrichment for genes associated with cytokine signalling in the immune system (normalised enrichment score = 2.22) (Fig. 3). In addition, GSEA identified a high over-representation of pathways involved in innate and adaptive immune response, within our differentially expressed genes, showing a massive activation of immune response in the HZn group compared with the MZn group after 1 week of feeding the zinc oxide diets.
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Mid- and long-term effects of feeding zinc oxide

After 2 weeks of feeding the different Zn diets, the HZn group had higher transcript amounts of FOXP3, the master transcription factor of regulatory T-cells\(^{27}\), in the HZn group compared with the MZn group \((P<0.01)\) and LZn \((P<0.05)\) groups within the GALT (Fig. 4(a)). In particular, we observed a higher expression of FOXP3 in JELN \((P<0.05)\) and JEPP \((P<0.05)\), as well as in ILLN \((P<0.05)\), in the HZn feeding group when compared to the LZn and MZn groups after 2 weeks of feeding the Zn diets (Fig. 4(b)).

We confirmed increased frequencies of CD4\(^+\)CD25\(^{high}\) regulatory T-cells among ILLN and JEPP and ILPP, respectively, in the HZn group compared with the MZn group \((P<0.05)\) by phenotypic analysis of immune cells using FCM (Fig. 4(c)) and (d)). Moreover, JELN and PAPIL showed comparable tendencies in the same direction, which is a higher relative cell count of CD4\(^+\)CD25\(^{high}\) regulatory T-cells among ILLN and JEPP and ILPP, respectively, in the HZn group compared with the MZn Zn feeding group \((P<0.01)\) after 4 weeks of feeding the Zn diets. The same tendencies could be seen in ileal lymph nodes (data not shown). In addition, we detected a lower relative transcript amount of TBET within GALT (Fig. 2(c)).

As expected, a significantly higher Zn concentration was detected in MLN of the HZn feeding group compared with the MZn group after 1 week \((P<0.05)\) of feeding. After 4 weeks, no difference in Zn concentration was observed (Fig. 1). Zn concentrations in MLN were surprisingly low compared with other tissues (jejunum, liver and pancreas)\(^{17}\). The low total Zn concentration in MLN, as well as the marginal changes in this concentration after feeding high levels of zinc oxide, indicates that MLN tightly control intra-cellular Zn levels. However, as no differential expression of Zn transporters was detected, the mode of regulation remains elusive.

Investigation of time-dependent, Zn-associated effects on adaptive immune cells upon HZn feeding showed that short-term feeding (1 week) led to significantly higher numbers of

### Discussion

Our result shows that immune cells derived from MLN and Peyer’s patches showed expression of ZIP4, ZnT1, ZnT2 and ZnT5 in all feeding regimens. To our knowledge, there is currently no information available that shows expression of Zn transporters in porcine immune cells. There is vast literature available linking Zn with immune function, as shown in the review paper by Wessels et al.\(^{28}\). Our hypothesis was that Zn transporters on the surface of immune cells respond to differences in Zn concentration. To validate this hypothesis, mRNA expression levels of the most common porcine Zn transporters (ZIP4, ZnT1, ZnT2 and ZnT5) were analysed. These genes were selected on the basis of previous research\(^{29,30}\), which showed differential expression of these transporters in epithelial cells and pancreatic cancer cells. However, in our current study, no differential expression of the selected Zn transporters was detected (Table 1).
activated T-helper cells (CD4+ and CD8αdim) ($P<0.01$). Molecular analysis revealed higher transcript amounts of IFNγ and TBET in the HZn group compared with the LZn group ($P<0.05$). TBET is the master transcription factor of T-helper 1 cells, whereas IFNγ is considered the main inducer of T-helper 1 cells (31). Our work provides support for the hypothesis that the differentiation of naive T-helper cells towards T-helper 1 cells might be short-term inducible by high levels of dietary Zn. In addition, a tendency was detected showing a higher relative cell count of cytotoxic T-cells within the HZn group compared with the MZn group after 1 week ($P<0.1$, data not shown).

GSEA on differentially expressed genes between MZn and HZn groups showed an over-representation of genes belonging to the pathway 'cytokine signalling in immune system'. This points towards an activation of the immune system after a 1-week feeding of zinc oxide.

To investigate this effect further, in vitro experiments can be used as a model to investigate direct effects of Zn on the immune cells. These in vitro experiments in which porcine MLN were treated with zinc acetate (Fig. 2), as well as zinc

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**Fig. 4.** (a and b) Relative mRNA expression of forkhead box P3 (FOXP3) in gut-associated lymphatic tissue (GALT) and in jejunal and ileocele mesenteric lymph nodes (JELN and ILLN, respectively), Peyer's patches from the jejunum and the ileum (JEPP and ILPP, respectively) and ileal papilla (PAPIL) analysed separately. (c) Exemplary original flow plot showing used gates to analyse cell populations. Plots are done using FlowJo version 7.9. displaying CD8-phycocerythin (PE) against CD4-fluorescein isothiocyanate (FITC) staining and CD25-allophycocyanin (APC) against CD4-FITC. (d) Cell counts, relative to the living lymphocyte population of CD4+CD25high regulatory T-helper cells shown for the same tissues as in (b). Differences were tested using a pairwise Mann–Whitney U test: * 0.01 < $P$ < 0.05, ** 0.001 < $P$ < 0.01, † 0.05 < $P$ < 0.1. Low zinc concentration (57 parts per million (ppm)); medium zinc concentration (164 ppm) and high zinc concentration (2425 ppm).

**Fig. 5.** Cell counts relative to the living lymphocyte population of CD4+ T-helper cells and CD8+ cytotoxic T-cells of jejunal mesenteric lymph nodes from piglets fed low zinc (57 mg zinc/kg) in diet (LZn), 164 mg zinc/kg (MZn) or 2425 mg zinc/kg (HZn) as zinc oxide for 4 weeks. Differences were tested using a pairwise Mann–Whitney U test: * 0.01 < $P$ < 0.05, † 0.05 < $P$ < 0.1.
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9. Pieper R, Vahjen W, Neumann K, et al. (2012) Dose-dependent effects of dietary zinc oxide on bacterial response and could provide an alternative to reduce the incidence of post-weaning diarrhoea in piglet. However, our findings indicate that long-term feeding (2–4 weeks) of high levels of Zn seems to have an immune-suppressive effect in post-weaned piglets and as such could potentially negate the benefits of Zn supplements in post-weaned piglets.

Acknowledgements

The study was funded by the German Research Foundation (Deutsche Forschungsgemeinschaft, DFG) within the Collaborative Research Group (SFB, Sonderforschungsbereich) 852/1 ’Nutrition and intestinal microbiota – host interactions in the pig’. The authors are solely responsible for the data and do not represent any opinion of neither the DFG nor other public or commercial entity.

S. K.-R., J. N. S. and D. K. performed the experiments; D. A., P. K. and S. K.-R. analysed the data; R. P., F. M. and V. G. contributed reagents/materials/analysis tools; J. Z. and G. A. B. conceived and designed the experiments and S. K.-R. and D. A. wrote the paper.

The authors declare no conflict of interest. The founding sponsors had no role in the design of the study; in the collection, analyses or interpretation of data; in the writing of the manuscript; and in the decision to publish the results.

Supplementary material

For supplementary material/s referred to in this article, please visit https://doi.org/10.1017/S0007114518002908

References


Weaning has an immune-activating effect on the adaptive T-cell response from an immune activation towards suppression. Higher relative cell counts of immune-suppressive CD4+CD25high regulatory T-helper cells were detected in all five analysed GALT. This was accompanied with higher transcript amounts of the FOXP3 gene, the master transcription factor of regulatory T-helper cells in the HZn-group compared to the MZn-group. Regulatory T-helper cells have a suppressive function on the inflammatory immune response and are reported to be capable of suppressing proliferation and differentiation of effector T-cells. As suggested by Pandiyan et al., this could be owing to induced Bcl-2-like protein 11-mediated apoptosis of activated effector CD4+ T-cells through the uptake of IL-2. In line with previous result, our results showed a reduction in the relative CD4+ T-cell count and also a tendency towards reduced relative cell counts of cytotoxic T-cells after 4 weeks in the HZn group compared with the MZn group. This is probably owing to the higher levels of regulatory T-helper cells detected after 2 weeks of feeding an HZn diet.

Zn taken orally is considered to be relatively nontoxic. However, toxicity symptoms have been known to occur with high levels of Zn intake in humans (approximately 100–300 mg Zn/d) when Zn is taken for longer periods of time (>10 months). Long-term Zn toxicity is often accompanied by a severe Cu deficiency (hypocupraemia); after cessation of the use of Zn supplements, the Cu levels often normalise. It is believed that the excess of Zn blocks the intestinal uptake of Cu, leading to hypocupraemia, until the excess of Zn is eliminated from the system. High levels of Zn intake (300 mg Zn/d) in humans for longer periods of time (6 weeks) have also shown to depress indices of immune function in healthy adults compared with baseline levels before supplementation. However, this effect was not seen in an elderly population exposed to lower levels of supplementation (100 mg Zn/d).

In a previous study with a similar experimental setup, we observed after 6 weeks feeding of HZn levels (2500 mg Zn/kg as zinc oxide) a decreased relative cell count of CD2+ T and natural killer cells, CD8+CD25high cytotoxic T-cells and CD4+ T-helper cells in MLN comparable to the situation in the LZN feeding group. This indicates a T-cell suppressive effect for both Zn deficiency and Zn excess. Interestingly, weight gain in the HZn group was only observed during the first week, but the weight gain was lower by the third week of the experiment compared with optimal- and low-level Zn diets, suggesting an optimal feeding time of 2 weeks when feeding high levels of zinc oxide.

In summary, our research suggests that short-term feeding (1–2 weeks) of high levels of Zn at the critical time point of weaning has an immune-activating effect on the adaptive T-cell response and could provide an alternative to reduce the incidence of post-weaning diarrhoea in pig handrory. However, our findings indicate that long-term feeding (2–4 weeks) of high levels of Zn seems to have an immune-suppressive effect in post-weaned piglets and as such could potentially negate the benefits of Zn supplements in post-weaned piglets.