Introducing enteral feeding induces intestinal subclinical inflammation and respective chromatin changes in preterm pigs

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Aim: To analyze how enteral food introduction affects intestinal gene regulation and chromatin structure in preterm pigs. 

Materials & methods: Preterm pigs were fed parenteral nutrition plus/minus slowly increasing volumes of enteral nutrition. Intestinal gene-expression and chromatin structure were analyzed 5 days after birth.

Results: Enteral feeding led to differential upregulation of inflammatory and pattern recognition receptor genes, including \( IL8 \) (median: 5.8, 95% CI: 3.9–7.8 for formula; median: 2.2, 95% CI: 1.3–3.3 for colostrum) and \( TLR4 \) (median: 3.7, 95% CI: 2.6–4.8 for formula; no significant differences for colostrum) with corresponding decondensed chromatin configurations. On histology this correlated with mild mucosal lesions, particularly in formula-fed pigs. In CaCo-2 cells, histone hyperacetylation led to a marked increase in \( TLR4 \) mRNA and increased \( IL8 \) expression upon stimulation with lipopolysaccharide (median: 7.0; interquartile range: 5.63–8.85) compared with naive cells (median: 4.2; interquartile range: 2.45–6.33; \( p = 0.03 \)).

Conclusion: Enteral feeding, particular with formula, induces subclinical inflammation in the premature intestine and more open chromatin structure in key inflammatory genes. This may increase the susceptibility for necrotizing enterocolitis.

Keywords: chromatin • DNA methylation • enteral nutrition • necrotizing enterocolitis • premature infants
In NEC development both in humans and pigs, TLR4 mRNA was not upregulated in preterm versus term pigs [17]. A possible explanation may be that TLR4 expression is modulated by environmental factors, such as feeding and bacterial colonization. Theoretically, these might induce changes in chromatin structure and CpG signaling, possibly destabilizing the intestinal homeostasis and thus creating the basis for an inappropriate immune response upon further exposure to bacterial antigens. Postnatal, introduction of enteral nutrition and bacterial colonization represent the most important environmental factors for the GI tract also playing a key role in NEC development both in humans [18] and pigs [19]. Importantly, early but slow introduction of enteral feeding using mother’s milk or colostrum is associated with significantly less NEC, relative to formula.

Against this background, we aimed to provide new insights into how introduction of enteral nutrition influences the transcriptional regulation in the intestine and how this contributes to the development of NEC. We analyzed the in-depth intestinal responses to a gentle enteral nutrition (EN) regimen as currently used in many European neonatal intensive care units with early and slow introduction of enteral feeds in association with supportive parenteral feeding. Therefore, we used preterm pigs since they tend to develop NEC spontaneously if fed larger amounts of formula particularly if feeding is advanced rapidly after preterm birth, closely reflecting human disease [6, 19–20]. This unique approach enabled us to analyze digestion- and inflammation-related genes, together with their DNA methylation and chromatin structure in the intestinal epithelium prior to development of clinical symptoms of NEC.

Materials & methods

Porcine model, enteral feeding & tissue collection

Fifteen preterm pigs (landrace × large white × duroc) were delivered by caesarean section on day 105 (90%) of gestation from one sow as previously described [19] and stratified according to birth weight and sex. Due to species-dependent organ maturation, the intestine matures relatively late in pigs and a preterm pig intestine at 90% gestation therefore represent the preterm human infant intestine at approximately 70% gestation [19, 21]. Housing in incubators with regulated temperature and oxygen supply, umbilical and orogastric catheterization and passive immunization were carried out as previously described [22].

The pigs were randomly assigned to either receive total parenteral nutrition (TPN) via the catheter fited in the umbilical artery (n = 5) or minimal enteral nutrition with bovine colostrum (n = 5) or preterm infant formula (n = 5). Bovine colostrum was used since this diet has been shown to elicit similar responses as porcine colostrum in the premature intestine of pigs [22] and is currently tested for use in human premature infants during the first days after birth [23]. All animals were also provided with supplementary parenteral nutrition (PN) to match the energy intake of the TPN group. The PN solution (Kabiven, Fresenius Kabi, Bad Homburg, Germany) for all groups was prepared and administered as previously described and modified to meet the nutrient requirements of preterm pigs [20]. TPN pigs received PN at a rate of 3, 4, 5 and 6 ml/kg bodyweight/h on days 1, 2, 3 and 4, respectively. The enteral feeding volume was 2, 4, 6, 8 ml/kg bodyweight/3 h on days 1–4. Formula adopted was a 16.5% solution of enfalac premature powder (Mead Johnson Nutrition, IL, USA) in bottled water and the colostrum feeding was a 17% solution of bovine colostrum powder (Biofiber-Damino, Gesten, Denmark). All procedures were approved by the National Council on Animal Experimentation in Denmark (protocol number 2012-15-293400193).

Pigs were euthanized and the GI tract (GIT) was sampled on the fourth day after initiation of EN. Clinical signs of NEC were recorded according to our macroscopic NEC evaluation system as described previously [19]. Three regions of small intestine, stomach and colon were graded for pathology using a macroscopic NEC lesion score (1: no or minimal focal hyperemic enterocolitis; 2: mild focal enterocolitis; 3: moderate locally extensive enterocolitis; 4: severe focal enterocolitis; 5: severe locally extensive hemorrhagic and necrotic enterocolitis or 6: severe extensive hemorrhagic and necrotic enterocolitis). Pigs with a score of 3 or more in any of the intestinal regions (Prox, Mid, Dist or Colon) were classified as a case of NEC (Supplementary Data 1; see online at www.futuremedicine.com/doi/full/10.2217/EPI.15.13). One to two 1 cm sections of the middle small intestine were stored in 4% paraformaldehyde for later histological analysis, two sections were snap frozen in liquid nitrogen and stored at −80°C for later intestinal brush border enzymatic activity assays of lactase, aminopeptidase A (ApA) and dipeptidyl peptidase 4, RT-qPCR and epigenetic analysis, respectively.

We decided to primarily focus our analyses on the mid intestine for two main reasons: The small feeding volumes used in this study were expected to have more immediate effects on the proximal and middle part of the intestine when compared with the lower...
parts; in preterm pigs the mucosal immune system including Peyer’s patches are not clearly evident in the distal intestine but are well developed in the mid intestine which thus probably best represents the premature human intestine regarding structural, functional and immunological responses.

Oligonucleotides
Primers utilized in this study are listed in Supplementary Data 2.

Sample pretreatment
Porcine intestinal specimens stored in liquid nitrogen were pulverized prior to the purification of RNA, gDNA and chromatin using a stainless steel mortar and piston (Cellcrusher, Cork, Ireland) prechilled in liquid nitrogen. Only tissue samples without microscopic signs of leukocyte invasion were used for further analysis.

High-throughput sequencing of the gut microbiota
The fecal microbiota of formula and colostrum-fed pigs was determined using tag-encoded 16S rRNA gene MiSeq-based (Illumina, CA, USA) pair-ended high-throughput sequencing. DNA storage condition, cellular RNA extraction and sequencing library preparation steps were conducted as previously described [24]. Quantitative Insight Into Microbial Ecology open source package (1.7.0) and CLC Genomic Workbench 7 (CLC bio, Aarhus, Denmark) were used to analyze the sequencing data as previously described [24]. The relative distribution of the genera registered was calculated for unified (85% of the most indigent sample) and summarized in the family and genus level OTU tables. Two samples were discarded due to the low number of reads. The alpha diversity measures expressed with an observed species (sequence similarity 97% OTUs) value were computed for rarefied OTU tables (28,000) using the alpha rarefaction workflow. Differences in alpha diversity were determined using a t-test-based approach employing the nonparametric (Monte Carlo) method (999 permutations) implemented in the compare alpha diversity workflow. The ANOVA test performed based on 999 rarefied OTU tables was used to determine quantitative (relative abundance) association of genus level OTUs with one of the two categories.

Gene-expression analyses
Prior to gene-expression analyses RNA was isolated from at least three biological replicates using Trizol reagent (Sigma-Aldrich, MO, USA) upon manufacturer’s recommendations. RNA integrity was validated utilizing an Agilent Bioanalyzer 2100 with the Agilent RNA 6000 Nano Kit for microcapillary electrophoresis. Using the RT2 First Strand kit (Qiagen/SABiosciences, Venlo, The Netherlands), cDNA templates were synthesized from 1μg RNA. Gene-expression was analyzed from these cDNAs using primer pairs as listed in Supplementary Data 2 with Quantitect SYBR Green Taq polymerase mastermix (Qiagen) on a Corbett Rotor-Gene 6000 qPCR device. PCR conditions were as follows: 95°C for 15 min, 40 cycles of (95°C for 15 s, 60°C for 30 s). Melting of PCR product was performed using a temperature gradient from 55–95°C, rising in 0.5°C increments. For relative comparative quantification of gene-expression fold changes we utilized the AΔCt method [25] using at least three housekeeping genes for normalization.

DNA CpG methylation analyses
For the assessment of DNA methylation in the promoter regions of differentially regulated genes, we combined CpG methylation-sensitive restriction endonucleolytic digests using HpaII (New England Biolabs, MA, USA) in combination with quantitative real time PCR. In addition, methylation-insensitive MspI (New England Biolabs) was used to control the integrity of the restriction enzyme recognition motif in each sample. Three sites within the H19 promoter were selected as DNA methylation-positive controls. Genomic imprinting of the H19 gene involves DNA methylation which leads to subsequent DNA hemimethylation [26]. Restriction digests were incubated at 37°C for 1 h prior to heat inactivation for 15 min at 95°C, simultaneously activating hot start Taq polymerase. PCR conditions were as described above for qPCR gene-expression analysis. Finally, melting of PCR product was performed using a temperature gradient from 55–95°C, stepwise rising in 0.5°C increments. For relative comparative quantification of template recovery after methylation sensitive restriction digests we applied a variation of the ΔΔCt method to compare the relative ratio of HpaII-digested DNA with undigested template DNA:

- \( \Delta \Delta Ct = Ct_{\text{HpaII-digested DNA}} - Ct_{\text{undigested DNA}} \)
- DNA recovery [%] = AE\(\Delta C_i\) x 100%, whereby AE is the amplification efficiency determined for each primer pair;
- Methylated CpG [%] = 100% - DNA recovery [%].

Benzonase endonuclease hypersensitivity assay
To analyze chromatin structure at specific sites of interest we applied the tissue accessible chromatin method [27] in combination with quantitative real time PCR, since Benzonase endonuclease hypersensitive sites frequently correlate with actively transcribed genes and are markers of cis-regulatory DNA elements such
as promoters, enhancers and locus control regions [28]. Briefly, liquid nitrogen-chilled, pulverized tissue samples were treated with the DNA endonuclease benzonase (Sigma-Aldrich). Next, DNA was purified by phenol: chloroform: isooamylc alcohol (25:24:1) extraction and precipitated with ethanol. Recovery of template DNA was compared with nondigested DNA analyzed using quantitative PCR as described above.

**Trichostatin A & lipopolysaccharides stimulation in CaCo-2**

Human CaCo-2 cells were maintained in Dulbecco’s Modified Eagle’s Medium enriched with 10% fetal bovine serum, 10% L-glutamine, 50 units/ml penicillin and 50 μg/ml streptomycin in a humidified atmosphere (5% CO₂) at 37°C. The medium was supplemented with 330 nM trichostatin A (TSA) (Sigma-Aldrich) and/or 50 ng/ml lipopolysaccharides (LPS) from Salmonella enterica serotype typhimurium (Sigma-Aldrich) as follows: treatment with TSA only for 1 h; treatment with LPS only for 2 h or treatment with TSA for 1 h followed by LPS exposure for 2 h. RNA-isolation, integrity testing and cDNA synthesis were performed as described above. PCR conditions were as follows: 95°C for 15 min, 50 cycles of (95°C for 15 s, 60°C for 30 s). Melting point analyses were performed utilizing qPCR as described above. All utilized primers are listed in Supplementary Data 2. The mitochondrial gene ATP 6 was used for normalization as it has no chromatin organization.

**Statistical analysis**

mRNA quantification levels of all analyses were obtained from triplicate real-time PCR measurements. Data are presented as medians ± interquartile ranges (IQR), minima and maxima. Testing for significant differences between groups was performed using the one-way ANOVA test for values with Gaussian distribution and the Mann–Whitney test for values without Gaussian distribution. When significant differences were found, the post hoc Bonferroni correction was applied to correct for use of multiple comparisons. The Kolmogorov–Smirnov test was utilized to rule out non-Gaussian distribution. In addition, one-way ANOVA testing was always combined with Bartlett’s test of equal variance. Values of p < 0.05 were considered statistically significant. All analyses were performed with GraphPad version 5.01 (CA, USA).

**Results**

**Enteral nutrition induces subclinical, diet-dependent intestinal epithelial inflammation**

When animals were sacrificed on day 5, none of them had any clinical symptoms of NEC (vomiting, abdominal distention, apnea, discoloration, bloody stools). Four out of the five formula-fed pigs but none of the colostrum-fed pigs showed clear NEC-like lesions in the colon region (NEC score 4 or 5) and were thus diagnosed with NEC despite the apparent absence of clinical symptoms prior to euthanasia. Characteristics of the pigs are given in Supplementary Data 1. For all pigs, histological examination of the mid small intestine revealed an intact intestinal epithelium with only mild mucosal inflammation and without any obvious NEC-like lesions in any of the three groups, although pigs fed with enteral diets had the most dense mucosal epithelium and longest villi (Figure 1A). This was also reflected by a significantly higher intestinal weight (Figure 2A), and intestinal weight per length (84 vs 68 mg/cm, p < 0.05) in both groups fed supplemental EN, relative to TPN. Lactase activity was markedly reduced in the pigs supplemented with formula (Figure 2B), while both groups receiving enteral nutrition showed elevated activity of dipeptidyl peptidase 4 (Figure 2C, p = 0.08) and ApA (Figure 2D), relative to TPN.

We also analyzed the composition of the microbial flora in the colon luminal content comparing formula with colostrum fed pigs. The sequences purged from chimeric reads yielded 673,377 giving an average of 84,172 sequences per sample (min = 33,171; max = 125,526; SD = 29,416 with a mean sequence length of 439 bp (SD = 12 bp). Alpha diversity measures expressed with observed species index showed no differences between the two categories (data not shown). The majority (>99%) of all reads were distributed between phyla Firmicutes and Proteobacteria in both categories accounting respectively for 43 and 57% in the formula-fed group, and 7 and 92% in the colostrum-fed group. These differences however did not reach significance. The genus level analysis indicated differences between the two categories reflected in the relative abundance of genus Klebsiella (proteobacteria), Enterococcus (firmicutes) and unclassified genus from Enterobacteriaceae family (proteobacteria). However, only the latter difference has reached significance (ANOVA p < 0.001***) (Figure 3).

**Nutrition induces diet-specific changes in gene-expression & chromatin signatures of genes related to inflammation & innate immunity**

The expression profiles of several genes differed significantly between TPN and supplemental enteral nutrition in general and between formula and colostrum feeding in particular as determined by analyses of relative mRNA enrichment by quantitative real time PCR (Figure 4). Relative mRNA copy numbers of inflammatory and pattern recognition receptor (PRR) genes, such as *IL8, TLR2, TLR4, REG3A*, an antimicrobial
protein in control of bacterial proliferation [29] and LYZ, an enzyme catalyzing hydrolysis of bacterial cell wall components, were elevated in both groups, compared with TPN (Figure 4A & B). This effect was significantly more pronounced for most inflammatory and innate immunity genes in the formula versus colostrum pigs (Figure 4C), with the only exception being that of TLR2, which was less expressed in formula-fed pigs. Interestingly, mRNA encoding the anti-inflammatory cytokine IL10, which is known to block NF-κB activity and be involved in the regulation of the JAK-STAT signaling pathway, was significantly enriched in pigs fed colostrum and downregulated in formula-fed piglets when compared with pigs fed PN alone (Figure 4). To answer the question whether the observed changes in gene-expression were also associated with changes in the chromatin structure for specific genes of interest, we analyzed endonuclease hypersensitivity sites. Importantly, almost all inflammatory and innate immunity genes, previously found to be differentially upregulated in formula-fed pigs, colocalized with chromatin segments exhibiting significantly higher benzonase endonuclease hypersensitivity, particularly TLR4 and IL8. The only exception was TLR2 (Figure 5A). Moreover, the observed differences in benzonase endonuclease hypersensitivity also seemed to correlate well with similarly directed changes in DNA methylation profiles (Figure 5B). Again no differences were found at the TLR2 locus (Figure 5B) or at any of the sites examined for other genes of interest (data not shown).

Chromatin decompaction through impaired histone deacetylation increases TLR4 expression in intestinal epithelial cells & leads to elevated IL8 mRNA accumulation upon subsequent LPS stimulation.

Next, we analyzed whether an induced open chromatin conformation in intestinal epithelial cells, specifically at the TLR4 locus might influence the inflammatory response upon stimulation with LPS. This was done using CaCo-2 cells, since under specific conditions CaCo-2 cells functionally resemble enterocytes in the small intestine [30] and currently represent one of the best characterized in vitro models of the intestinal barrier [31]. CaCo-2 cells were treated with the histone deacetylase (HDAC) TSA and their inflammatory response was compared with naive CaCo-2 cells. Treatment with TSA resulted in significantly higher mRNA copy numbers for TLR4, whereas it had no detectable effect on TLR2 and IL8 mRNA (Figure 6). Stimulation with LPS led to a significant increase in IL8 mRNA copy numbers in both TSA pretreated and naive CaCo-2 cells, but the observed increase was significantly higher in TSA pretreated cells (median: 7.0; IQR: 5.63–8.85 vs 4.2; IQR: 2.45–6.33; p = 0.03) (Figure 6). No differences were noted for IL6 mRNA copy numbers, which were found to be decreased upon TSA treatment and/or upon LPS stimulation, when compared with naive CaCo-2 cells.

Discussion

The protective effect of breast milk toward the development of NEC was first reported several decades ago [32], but nevertheless it has remained unclear how the timing (early vs late), amount (slow vs rapid) and diet (breast milk vs formula) of the first enteral feeds for preterm neonates affect NEC sensitivity [33]. In this study, we used a preterm pig model to investigate the effect of an early and slow introduction of enteral feeds on the intestine as used in many neonatal intensive care units for preterm infants. This slow feeding advancement generally preserved small intestinal structure and functions and induced only mild NEC lesions (in the colon, and only in formula-fed pigs) in contrast to more rapid advancement which has been shown to induce clinical NEC symptoms and mac-
roscopic intestinal pathology in preterm pigs, especially using formula diets [34]. Thus, we were able to characterize the intestinal epithelial response on the gene-expression and chromatin level to the first enteral feeds prior development of any clinical signs of NEC. We also compared an infant formula with a bovine colostrum product that has previously been shown to protect against NEC in newborn preterm pigs. The results enabled us to provide representative data on how the first small amounts of enteral feeds can shape the epigenetic landscape and consequently influence the transcription of genes in the premature intestinal epithelium and furthermore, how these changes might influence the susceptibility for NEC development.

Similar to previous studies in preterm pigs, using rapid transition to higher amounts of enteral formula or colostrum (120 ml/kg/day) [20], we observed lowered intestinal mass and activity of brush border enzymes such as dipeptidyl peptidase 4 and ApA in TPN-fed pigs when compared with the enteral feeding groups. This most likely reflects lack of enteral stimulation of tissue metabolism and blood flow. Interestingly, lactase activity was specifically reduced in pigs fed small amounts of enteral formula, even in the absence of any NEC lesions in the small intestine. Probably, intestinal lactase activity was particularly sensitive to the suboptimal formula diet because this enzyme – in contrast to other brush border enzymes – is most abundantly expressed in the proximal and middle intestine by the oldest enterocytes on the villus tips that may first show the effects of an adverse diet response. This reduced lactase activity may facilitate bacterial overgrowth due to an increased intraluminal availability of undigested carbohydrates thus predisposing to later development of NEC [20,35].

More importantly, enteral feeds, and formula feeding in particular, induced subclinical inflammation in

Figure 2. Influence of diet on intestinal weight and digestive enzyme activities in preterm 5-day-old pigs. Preterm pigs were fed exclusively with TPN, parenteral nutrition plus formula (formula) or parenteral nutrition plus colostrum (colostrum). (A) Small intestinal weight and (B) enzymatic activities of lactase, (C) dipeptidyl peptidase 4 and (D) ApA differed significantly among the three groups (means ± standard error of mean). *p < 0.05; **p < 0.01.

TPN: Total parenteral nutrition.
the intestinal epithelium with significant upregulation of inflammatory genes, such as IL8 and PRR, such as TLR4. This translated to corresponding changes in epigenetic signatures. Thus, DNA of upregulated genes, such as IL8 and TLR4, was significantly more sensitive to benzonase, which preferentially digests DNA in regions of low nucleosome occupancy. Since such endonuclease hypersensitive regions directly reflect the accessibility of the genome [28], IL8 and TLR4, among others, seemed to be localized in decondensed, putatively activated chromatin regions, which might be enhanced depending on the type of enteral nutrition during the first days after preterm birth. Moreover, lower levels of CpG methylation were particularly observed for IL8 after formula feeding, providing additional indication for long lasting gene activation, since differential CpG signaling is frequently associated with long-term changes in the regulation of gene-expression [56]. Such a baseline activation of inflammatory and PRR genes fits well with the current concept of an exaggerated innate immune response as key factor for NEC [12,14–15,37–38], particularly since the observed nutritional effect also involves TLR4. Since to date, no genetic markers could be associated with NEC, this diet-dependent activation could provide at least a partial explanation for the exaggerated innate immune response observed in the mouse model [37]. These results also help to understand why TPN-fed preterm neonates are less sensitive to NEC, although continued TPN has many negative consequences for the immature intestine, including decreased mucosal growth and defense. In fact, in this study, TPN led to a decreased expression of antimicrobial peptides REG3A and LYZ. This most likely reflects Paneth cell dysfunction thus increasing the susceptibility to mucosal bacterial invasion, as reported for the mature intestine [39]. Possibly such Paneth cell dysfunction plays an important role in the pathogenesis of NEC as recently suggested by McElroy et al. [40]. In consequence, optimization of postnatal nutrition in the premature intestine is obviously more complicated, particularly since it still remains unclear whether early trophic feeding in human premature infants is beneficial or not [41].

Preterm pigs fed colostrum showed a less pronounced intestinal expression of inflammatory and PRR genes and a higher expression of the anti-inflammatory cytokine IL10, relative to pigs-fed formula or parenteral nutrition alone. Remarkably, IL10 can block NF-κB activity as an important regulator of immunity and inflammation, which can also be activated by TLR4 stimulation [42]. This finding of increased IL10 expression upon colostrum feeding might thus explain the previously reported reduced NF-κB activity in the presence of upregulated TLR4 in colostrum-fed pigs [43]. At the same time, it partly resolves the contradictory results between the mouse model showing a TLR4-mediated exaggerated innate immune response at the center of NEC pathogenesis [37] and the porcine data demonstrating higher TLR4 expression in NEC-protected colostrum-fed pigs [37,43]. Finally, it provides a reasonable explanation for the protective effect of colostrum in the premature pig model [39] and possibly for the protective effect of breast milk in humans [44]. At this point it is important to mention that we also observed some variations in the composition of colonic microbiota between colostrum and formula-fed pigs. Even though statistically not significant except for one genus, these variations might have partially contributed to the observed differences in gene-expression and chromatin conformation between colostrum and formula-fed pigs. However, this finding was surprising since in a similar study feeding with higher volumes of enteral feeds only showed marginal differences in bacterial colonization between formula- and colostrum-fed preterm pigs. However, this finding was surprising since in a similar study feeding with higher volumes of enteral feeds only showed marginal differences in bacterial colonization between formula- and colostrum-fed preterm pigs.

Figure 3. The average relative distribution of the major gut microbiota genera (>0.1% in both categories) determined for formula (n = 4) and colostrum-fed (n = 4) pigs. Taxa denoted as ‘unclassified’ means that the reference database does not have an official taxonomy for this cluster. Taxa denoted as ‘other’ (unidentified), indicates ambiguity in the assignment meaning that more than one bacterial taxa could be assigned to this cluster. An unclassified genus within the Enterobacteriaceae family was the only taxon significantly different between the two categories (ANOVA p < 0.001*).
A  

**Formula vs TPN**

- **Fold change**
  - REG3A
  - ESPE1
  - LYZ
  - IL10
  - TLR2
  - TLR4
  - H19
  - KRT8
  - RPL19
  - ACTB
  - PCK2
  - GAPDH
  - TUBB3a
  - DEFB1
  - ASRGL1
  - CRYL1

B  

**Colostrum vs TPN**

- **Fold change**
  - REG3A
  - ESPE1
  - LYZ
  - IL10
  - TLR2
  - TLR4
  - H19
  - KRT8
  - RPL19
  - ACTB
  - PCK2
  - GAPDH
  - TUBB3a
  - DEFB1
  - ASRGL1
  - CRYL1

C  

**Formula vs colostrum**

- **Fold change**
  - REG3A
  - ESPE1
  - LYZ
  - IL10
  - TLR2
  - TLR4
  - H19
  - KRT8
  - RPL19
  - ACTB
  - PCK2
  - GAPDH
  - TUBB3a
  - DEFB1
  - ASRGL1
  - CRYL1
pigs [45]. Moreover, bacterial colonization of the gastrointestinal mucosal surfaces in mammals begins at birth and does not initially appear to depend much on diet but rather on host-response mechanisms and the maternal microbiota in the first week of life [46,47]. Only later in life the relatively stable gut microbial composition may be more influenced by the type of enteral nutrition [48]. Thus the relevance of the observed variations in colonic microbiota in this study – particularly considering the low number of animals – currently remains unclear but warrants further evaluation.

In the last part of this study, we aimed to assess the possible influence of an open chromatin conformation on the TLR4-mediated innate immune response of the intestinal epithelium. We could show that inhibition of HDAC using TSA, inducing chromatin hyperacetylation and an open chromatin structure [49], led to increased TLR4 expression in CaCo-2 cells, whereas TLR2 expression remained unaffected. This is in accordance with our observations from the animal model. Strikingly, subsequent stimulation with LPS, the major component of Gram-negative bacteria that activates TLR4 [50] elicited a significantly higher inflammatory response as measured by relative IL6 mRNA copy numbers in TSA pretreated CaCo-2 cells. In contrast, IL6 expression remained almost unchanged upon stimulation with LPS which is in line with a previous report using the same cell line [51]. We also found a significant decrease in IL6 mRNA expression upon TSA treatment, confirming that TSA treatment has no effect on IL6 promoter accessibility as previously reported [52]. Even though speculative, the observed decreased IL6 mRNA levels might thus result from an upregulation of counteracting cytokines, such as IL10, known to be upregulated in response to HDAC inhibition [53].

Limitations of this study
The most important limitation of this study is the lack of data from preterm infants. Even though preterm pigs and infants share many physiological and intestinal similarities, the immature porcine gut and immune system (innate as well as adaptive) still differs from those in preterm infants. Consequently, larger studies, preferably using infant intestinal samples, are necessary to confirm the findings from this study. Also, to prove that the observed intestinal responses to enteral nutrition eventually lead to NEC, it is necessary to replicate the findings in a larger group of animals followed for a longer time with serial intestinal samples for each individual. Finally, we only analyzed the microbiota from the colon luminal content and not from the mid intestine. Even though it is reasonable to assume that differences in mid intestine microbiota translate to colonic microbiota, the exact role of the microbiota for the observed effects need further study. Nevertheless, we are convinced that optimization of the enteral nutrition regimen in the days and weeks just after preterm birth is a key tool to reduce the high susceptibility to NEC.

Conclusion
In summary, here we provide evidence that enteral nutrition might at least partially be the basis for the observed exaggerated innate immune response seen in NEC. We demonstrated that even a gentle enteral feeding regimen with a gradual increase in enteral feeds as used by many neonatal intensive care units induces subclinical inflammation in the premature intestinal epithelium with subsequent changes in chromatin conformation in inflammatory and PRR genes. Importantly, these changes occurred in the absence of any clinical symptoms suggestive of NEC which rarely develop until preterm pigs or infants approach full enteral feeding (e.g., 120–180 ml/kg/day). Also, this inflammatory response was much more pronounced using infant formula, as first diet, compared with bovine colostrum.

Future perspective
We have recently initiated preliminary studies on preterm infants using bovine colostrum as the first enteral diet, prior to the later transition to full feeding with formula or breast milk. Together with additional studies these will help to further characterize the influence of enteral nutrition in modulating intestinal gene-expression and chromatin structure. These data will then allow us to adapt enteral nutrition for premature infants possibly even in a personalized way thus reducing the risk for intestinal dysfunction and NEC in particular. For this process of personalized medicine a detailed genetic characterization of NEC susceptibility would be highly desirable.
Figure 5. Influence of diet on intestinal chromatin configuration and CpG methylation. Genes upregulated in formula (dark grey), compared with colostrum (light grey) piglets, such as IL8, TLR4, REG3A and LYZ, were associated with a significantly more open chromatin conformation. (A) TPN-fed piglets were used as reference the groups supplemented with either formula or bovine colostrum. (B) CpG methylation patterns also correlated with mRNA levels. The IL8 locus was significantly less methylated in pigs fed colostrum (co) and particularly in pigs-fed formula (fo) when compared with pigs fed (PN) alone (iv). The only exception was TLR2. Methylation levels at the H19 locus were about 50% in all cases reflecting its status as imprinted gene. Data are presented as median ± interquartile ranges, minimum and maximum. All experiments were performed in technical triplicates. TPN: Total parenteral nutrition, PN: parenteral nutrition.
Figure 6. Inflammatory response upon lipopolysaccharides stimulation in CaCo-2 cells with or without treatment with trichostatin A. Upon histone deacetylase inhibition using TSA in CaCo-2 cells, TLR4 mRNA levels increased 12.7-fold whereas TLR2 and IL8 mRNA levels remained unchanged. Stimulation with LPS in naive CaCo-2 cells induced an increase in IL8 mRNA levels. This effect was significantly more pronounced in TSA pretreated CaCo-2 cells. Data are presented as the median ± interquartile range, minimum and maximum. All experiments were performed in technical triplicates. Asterisks indicate p < 0.05. LPS: lipopolysaccharides; TSA: Trichostatin A.

reviewed the manuscript. A Jenke, P Sangild and J Postberg took part in the main study design, data analyses and supervision of the study. L Krych was responsible for the microbiota analyses. A Jenke had primary responsibility for final content. All authors read and approved the final manuscript.

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Executive summary
- Enteral nutrition induces subclinical intestinal epithelial inflammation
- Histopathological inflammation is more pronounced using formula feeding.
- Lactase activity was markedly reduced in the pigs supplemented with formula.
- Nutrition induces diet-specific changes in gene expression and chromatin signatures of genes related to inflammation and innate immunity
- Upregulation of several inflammatory and pattern recognition receptor genes correlates with histopathology and is more pronounced in formula feeding.
- Only colostrum induces expression of IL10 possibly lowering necrotizing enterocolitis susceptibility.
- Changes in gene expression translate to diet-specific changes in chromatin conformation.
- Chromatin decompaction increases Toll-like receptor 4 expression and lipopolysaccharides responsiveness in vitro.
est in or financial conflict with the subject matter or materials discussed in the manuscript apart from those disclosed.

No writing assistance was utilized in the production of this manuscript.

Ethical conduct of research

This study was conducted with approval of the Witten/Herdecke University Ethics board. All animal procedures were approved by the National Council on Animal Experimentation in Denmark (protocol number 2012-15-293400193). The authors state that they have obtained appropriate institutional review board approval or have followed the principles outlined in the Declaration of Helsinki for all human or animal experimental investigations. In addition, for investigations involving human subjects, informed consent has been obtained from the participants involved.

References

Papers of special note have been highlighted as:

- of interest; • of considerable interest.


•• This review provides a comprehensive summary on necrotizing enterocolitis (NEC) from the clinical perspective.


•• This excellent study provides the first data how breast milk might possibly alleviate NEC in the rodent model.


• The authors provide the first genome-wide association study (GWAS) data on human intestinal tissue comparing NEC with spontaneous intestinal perforation.


•• The study describes the currently best animal model for NEC.


22 Bjornvad CR, Thymann T, Deutz NE et al. Enteral feeding induces diet-dependent mucosal dysfunction, bacterial...
• This review summarizes the current evidence for a Toll-like receptor 4-driven exaggerated immune response as key factor for the development of NEC.


