DNA methylation as a regulator of intestinal gene expression

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Abstract
The intestinal tract is the entry gate for nutrients and symbiotic organisms, being in constant contact with the external environment. DNA methylation is one of the keys to how environmental conditions, diet and nutritional status included, shape functionality in the gut and systemically. This review aims to summarise findings on the importance of methylation to gut development, differentiation and function. Evidence to date on how external factors such as diet, dietary supplements, nutritional status and microbiota modifications modulate intestinal function through DNA methylation is also presented.

Key words: DNA methylation; Intestines; Nutritional status; Diet

Epigenetics refers to the regulation of gene expression through processes that do not change DNA sequence. These changes usually occur in response to environmental conditions and are inheritable through cell division. DNA methylation is the most studied and most common epigenetic modification, influencing gene expression when reduced in promoter regions, the binding site of enzymes responsible for transcription, and increased in the body and 3′ end of genes. In vertebrates, this is represented by a methyl group covalently bound to the carbon at the 5′ position of the pyrimidine ring of 5-methylcytosine (5mC) in a CpG dinucleotide (a cytosine followed by a guanine). Genomes CpG density is low, with the majority of CpG being present in CpG islands (CGI) that are on average 1000 bp long and have a relatively high concentration of hypomethylated CpG dinucleotides.

Approximately 70% of annotated promoters are associated with CGI, including promoters of tissue-specific genes and developmental regulators, with around 50% of the CGI in mice and humans located in transcription start sites. While the functionality of dynamic methylation patterns of the CGI has been more associated with repression of transposons (DNA sequences capable of moving from one location to another within the genome) and repetitive elements, imprinting (selective monodispersed expression), X-inactivation (inactivation of one copy of the X chromosome in females) and promoter accessibility to transcription factors, the fact that a difference in the methylation pattern reflects modifications of gene expression levels indicates that DNA methylation could be a marker for modification of transcriptional patterns due to environmental changes such as diet and microbiota composition. This review highlights difficult to create a causation link between DNA methylation, transcriptional activation and gene expression. However, most CGI promoters are protected against de novo methylation indicating a functional reason for its maintenance in a hypomethylated state. This is corroborated by the embryonic lethality of the knockout (KO) of the DNA methyltransferase (DNMT) genes Dnmt1 or Dnmt3h in mice and the early death of Dnmt3a-KO mice. The DNMT enzymes are responsible for the establishment and maintenance of methylation patterns on DNA while ten-eleven translocation (TET) enzymes are linked with DNA methylation regulation by catalysing the conversion of 5mC into 5-hydroxymethylcytosine (5hmC), which is the first step of the demethylation process.

In somatic cells, DNA methylation patterns are maintained through inheritance across mitoses with remarkable precision and can even be maintained across generations. However, the maintenance of methylation status is not universal in all genomic regions, as some regions without apparent regulatory functions are heterogeneously methylated even in cell clones.

The binding of TCF proteins that regulate transcription, can reduce the methylation level of local and flanking CGI, making it difficult to prove causality between DNA methylation and gene expression. However, the fact that a difference in the methylation pattern reflects modifications of gene expression levels indicates that DNA methylation could be a marker for modification of transcriptional patterns due to environmental changes such as diet and microbiota composition. This review highlights

Abbreviations: BS, bisulphite; CGI, CpG islands; CpG, 5′-cytosine–phosphate–guanine; CV, conventional; DMR, differently methylated regions; DNMT, DNA methyltransferase; GF, germ-free; ISC, intestinal stem cells; KO, knockout; LINE1, long interspersed nuclear element 1; MD, methyl donor; MSP, methylation-specific PCR; TCF4, transcription factor 4; TET, ten-eleven translocation; TLR4, toll-like receptor 4; 5hmC; 5-hydroxymethylcytosine; 5mC, 5-methylcytosine.

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evidence on how nutritional status, diet and microbiota influence intestinal functionality through DNA methylation (Fig. 1), including methodologies used to measure DNA methylation in intestinal tissue, links between DNA methylation and intestinal development, differentiation and maturation and the influence of microbiome modulation on the intestinal methylome.

Methodologies for the analysis of DNA methylation

Multiple methodologies have been used for the analysis of DNA epimodifications. As genome-wide approaches have become more viable and feasible due to improvements in DNA sequencing technology and analysis, epigenetic modification methodologies have also evolved to high-throughput technologies with single-base resolution.

The most used approaches for methylation profiling target one or a few genomic loci. Identifying CpG methylation on a sequence of defined length is frequently done by methylation-specific PCR (MSP) [40] or pyrosequencing [47]. With either methodology, the DNA is first treated with bisulphite (BS) for the conversion of unmethylated cytosines into uracils and subsequently into thymines, which allows the identification of differently methylated loci [48]. MSP utilizes primers, small oligonucleotides complementary to the targeted region, that are specific for a methylated or unmethylated locus and the presence of a band determines the methylation status [41]. One drawback is that only the region of the primers can be truly analysed and a CpG rich region is necessary to create primers specific enough to differentiate between methylated and unmethylated DNA. In BS pyrosequencing, the amplicon sequence can be analysed, which makes it a more quantitative and robust technique than MSP. Pyrosequencing entails sequencing-by-synthesis using pyrophosphate release to identify the nucleotide added to the sequence and therefore can quantify the ratio between cytosines and thymines at a given CpG position. It has also been extensively used as a validation technique for high-throughput methods [19–25]. MS-based approaches can also be used to analyse the methylation level at targeted regions [26].

Global methylation quantification is the measure of the general level of DNA methylation without considering methylation changes at specific loci. Different techniques are used to investigate global methylation levels, such as pyrosequencing of Alu or long interspersed nuclear element 1 (LINE1) elements, luminometric methylation assay and HPLC. PCR-based methods estimate the methylation status of the most prevalent repetitive elements in the genome, Alu and LINE1, by pyrosequencing, where the cytosine:thymine ratio at a given CpG in LINE/Alu elements indicates the methylation status. Luminometric methylation assay uses the ratio of the digestion of unsensitive:sensitive isoschizomers to estimate the methylation status. Pyrosequencing LINE1 elements showed the smallest variation between samples and corresponded the best to results obtained by HPLC which is one of the most accurate methods to assess methylation [27]. However, caution must be used in extrapolating evidence of methylation changes in LINE1 elements to global changes in specific conditions [28]. High performance capillary electrophoresis and chromatography approaches separate methylated and unmethylated DNA by size, and both are quantified, but despite being considered the gold standard for methylation quantification, they usually require a large amount of DNA [29,30]. MS also has high sensitivity and requires lower amounts of starting material [31,32].

For genome-wide methylation analysis, although microarray-based methodologies have been used extensively, they require whole genome amplification, which can insert sequence bias, and previous knowledge of the sequence for probe synthesis, but are much more cost-effective than next-generation sequencing approaches [33]. This method has been progressively replaced by whole genome bisulphite sequencing (MethylC-seq) that now is considered the gold standard approach and can identify differently methylated regions (DMR) with single bp resolution. Whole genome bisulphite sequencing is done by sequencing of the whole genome after BS treatment and only one of the strands is analysed, although alternative protocols can analyse all four strands formed after the BS treatment [34].

The use of techniques to enrich samples with methylated cytosines or CpG-rich regions of the genome can be used together with genome-wide approaches such as microarrays or next-generation sequencing. The use of beads that have high affinity to 5mC to precipitate methylated regions (methylated DNA immunoprecipitation) of the genome [35] is one enrichment technique. Methyl binding domain proteins can also be used to enrich methylated genome regions and although it can be used to compare regions between samples, it cannot assess single nucleotide methylation status [36,37].

The use of methylation-sensitive discordant isoschizomers can be used in conjunction with microarrays and sequencing, in, for example, the HpaII tiny fragment enrichment by ligation-mediated PCR (HELP) assay and reduced representation bisulphite sequencing, respectively, usually with the intent to enrich CpG containing areas. The HELP assay [38] has also been used together with sequencing, and in this way it has similar CpG coverage to reduced representation bisulphite sequencing. HELP-tagging is a modification of the methyl-sensitive cut counting assay that uses the HpaII restriction enzyme, with a normalisation step using reads formed by non-methylation-sensitive MspI cleavage. This approach works for CpG rich sites, as well as more CpG depleted regions in the genome, and compares reads formed by MspI or HpaII digestion to estimate the methylation at each of these sites [39].

Reduced representation bisulfite sequencing [40–42] uses restriction enzymes that have their recognition site in CpG dinucleotides and is insensitive to methylation (usually MspI); therefore, each fragment will contain two CpG (one on each end). After digestion and size selection of the fragments, the samples are BS treated to convert unmethylated cytosines into thymines and then sequenced. Reduced representation bisulfite sequencing is one of the most popular methylation assay techniques because it can be used to analyse 80% of the CGI and 60% of the promoters [43]. But as the other BS-based methods, it cannot distinguish between methylcytosine and hydroxymethylcytosine. A more complete review on these and other methods for methylation analyses can be found at Tost and Gut [44].
The intestinal tract is the organ in mammals responsible for digestion of food for nutrient extraction and absorption and for excretion of digestion and metabolic waste. It is exposed to the external environment and is in constant and intimate contact with micro-organisms in a symbiotic manner.

The small intestine, where most nutrients are absorbed under neurological and endocrine regulation, is a hollow tube lined by a mucosal layer consisting of a single layer of epithelial cells that are the physical barrier between the commensal bacteria and the interior of the host body. These epithelial cells are a heterogeneous population and are being constantly renewed by the division of intestinal stem cells (ISC) located in the intestinal crypts. Other self-renewal cell populations, such as haematopoietic stem cells, have been shown to have progressive changes in DNA methylation in a cell-specific manner (45).

Intestinal development undergoes transitional stages before and after birth, with the postnatal period being crucial for proper intestine and immune system maturation in response to microbial colonisation, oral nutrition, weaning and nutrient availability (46–49). Differences in both gene expression and DNA methylation patterns in the intestines were observed during the transitions between fetal, suckling, weaning and adult life periods (46,50), and differences between intestinal sections in response to maturation were also observed. Regional identity between the different intestinal sections has also been linked to stable DNA methylation signatures observed in paediatric and adult human intestinal epithelial organoids, with fetal-derived organoids presenting dynamic methylation changes suggestive of in vitro maturation (51). Notably, most genes that are differentially expressed at the transcriptional level between the suckling and weaning periods stabilise expression levels after weaning, suggesting that early life transitional states can influence gene expression in adult intestines (46).

Cryp ts are responsible for cell proliferation and turnover rate of the intestines and go through maturation in the early postnatal period. Using Dnmt1 KO mice, Yu et al. observed a large variance in the methylome of colonic stem cells of intestinal crypts in mice during the suckling period, especially during maturation, the transition from fetal ISC to adult ISC, compared with differentiation, when the stem cell is turned into different types of epithelial cells (50). The methylation occurred mainly as hypermethylation in CGI associated with gene bodies and the 3’ end of genes. Dnmt1 KO mice presented severe intestinal abnormalities and an 80% mortality rate, indicating that DNA methylation is indeed important to intestinal maturation and that dysregulation of methylation patterns in the early postnatal period can lead to an immature intestinal tract. Furthermore, the reduced methylation at the 3’ end of genes in the KO mice was associated with reduced gene expression. Corroborating with the study of Yu et al., Forn et al., using amplification of inter methylated sites sequencing (AIMS-Seq), found only few...
differences (only 1.65% of the amplicons) between differentiated villi cells and ISC in the small intestines of mice with hypermethylation being the vast majority of the changes\(^5\).\(^5\).

Other studies have demonstrated the importance of DNMT1 and methylome establishment during the normal maturation and differentiation processes of ISC, confirming differences between methylomes during maturation and differentiation\(^5\).\(^4\),\(^5\).\(^5\). They observed a tendency for decreased methylation during differentiation and that DMR were enriched at CGI and CpG shores with the majority (61%) of DMR hypomethylated during differentiation being at intronic regions. A correlation between loss of methylation in promoter, 3' untranslated region and first intron and increased expression level was observed, occurring mainly in genes associated with small intestine metabolism.

The DMR that gain methylation through differentiation were associated with signalling pathways important to ISC function such as wingless and Int-1 signalling pathways (Wnt) and overlap with binding sites for transcriptional factors in ISC such as caudal type homeobox 2 (CDX2)\(^5\).\(^5\). Sheaffer suggests that hypomethylation coordinates the binding of TCF; however, Bestor et al.\(^5\),\(^6\),\(^7\) and others\(^5\),\(^6\),\(^7\) propose that the binding of transcriptional factors causes the hypomethylation on the motif regions. Kaaïj et al.\(^5\),\(^6\),\(^7\) also related the binding of a TCF with hypomethylation, in this case TCF4, which the author suggests contributes to DMR formation during differentiation of small intestine leucine-rich repeat-containing receptor 5 positive cells (Lgr5+ cells), a marker of ISC\(^5\).\(^8\). Comparing fetal and paediatric intestinal epithelial cells, Kraiczey et al.\(^5\) found that in human ileum and colonic cells, the potentially regulatory DMR frequently overlap transcription start sites and that these DMR were enriched in pathways involved with embryonic, tissue and intestine development. In regulatory DMR-associated genes, the expression was inversely related to promoter DNA methylation. Age was also inversely related to methylation of innate immunity genes and positively correlated with those genes’ expression levels.

Differences in methylation level in enhancers and promoters of genes related to ISC function were observed and although intestinal DNMT1 ablation in adult mice intestines can cause alternative methylation patterns in enhancers and delay the differentiation process (by extension of the cryptic zone), it seems to have milder effects compared with early life KO\(^5\),\(^4\),\(^5\),\(^5\),\(^8\). Perinatal ablation of DNMT1 caused loss of nascent villi due to hypomethylation and premature differentiation and apoptosis\(^5\) due to DNA damage and genomic instability in progenitor cells linked to hypomethylation. Corroborating this, genes associated with cell cycle were expressed at lower levels in the DNMT1/-/- ablated cells along with up-regulation of P21, a gene expressed due to DNA damage and associated with cell cycle arrest\(^5\).\(^4\). Interestingly, Kaaïj et al.\(^5\) could not find many DMR in transcription start sites due to differentiation of Lgr5+ inter villi epithelial cells in adult intestines. Instead, they observed that there was considerably stable methylation status between the two cell types, which could corroborate the fact that DNMT1 and de novo methylation is more important during development than differentiation, hence the milder effects of DNMT1 in adult mice and stabilisation of gene expression after weaning.

Elliott et al. investigated the milder effect of 5\(\text{mC}\) to 5\(\text{hmC}\) conversion in intestinal cells compared with differentiated cells and the opposite was true for the Tet2 and Tet3 genes. Using 5\(\text{mC}\) mutant mice, it was shown that the mutant mice were growth retarded and that the mutation caused significant lethality in the early postnatal period compared with wild-type littermates\(^5\).\(^4\). Also, in the
Intestinal tissue, villi height and the number of progenitor cells in the crypts were significantly reduced in the Tet1+/− mice, indicating that the Tet1 gene is important for cell proliferation. Organoid budding and size were also decreased in the cultures derived from Tet1+/− intestines. Overall, 5hmC enrichment at the Wnt target site is essential for gene expression of Wnt target genes in Lgr5+ cells and that the depletion of the Tet1 gene product decreased the level of 5hmC and therefore decreased the expression level of those genes. The conversion of the 5mC into 5hmC is an important regulatory mechanism for the expression of Wnt target genes during intestinal postnatal maturation. Kraiczky et al. observed that Tet1 gained methylation and reduced expression through development\(^{19}\).

A recent study investigating 5hmC observed dynamic changes during differentiation of mouse progenitor adult ISC into epithelium specialised cells\(^{65}\). Interestingly, 5hmC in progenitors did not correlate with transcripts levels while a positive 5hmC gene expression correlation was observed after differentiation. Furthermore, ~60% of the 5hmC were intragenic and 5% of the intergenic 5hmC were located with 5 kb of a transcription start site with the gain of 5hmC upon differentiation occurring mainly in intergenic regions. Gene ontology analysis of the genes containing significant changes in 5hmC was enriched for cell metabolism and cell–cell interaction while intergenic 5hmC was assigned to the closest gene and those were enriched for organ morphogenesis, cell signalling and DNA template processes.

Overall, the importance of DNA methylation on early life intestinal development is notable. The regulation of intestinal maturation is being driven at least in some part by DNA methylation with the differentiation process being less dependent on methylation modifications. Also, it is interesting to note the association between methylation level differences during the growth/ageing processes, mostly with hypomethylation either locally in the first case or globally in the later.

**Microbiota composition influence on intestinal DNA methylation and development and ‘functionality’**

Intestinal commensal bacteria colonisation in the postnatal period influences physiology, morphology and functionality of the intestinal tissue\(^{66,67}\) and can affect gene expression of the host, as shown in many studies comparing germ-free (GF) and conventionally raised animals\(^{68–72}\) and reviews on the topic\(^{66,73,74}\). The link between microbiota and intestinal renewal has also been found in species that have historically less evidence of DNA methylation such as Drosophila\(^{75}\), indicating that DNA methylation status may be an acquired regulation during evolution. Also, epigenetic reprogramming of host genes has been demonstrated during microbial and virus infections and has been associated with malignant progression of cancer\(^{76,77}\).

Microbiota effects on host physiology can be associated with bacterial metabolites, such as butyrate, a by-product predominantly from Firmicutes metabolism, that potentially influences host gene expression by histone chromatin modifications\(^{78}\). Fermentation of dietary fibre by bacterial populations in the colon produces butyrate that is the main source of energy to colonocytes. Also, the regulatory function of butyrate as an inhibitor of histone deacetylase is consequently associated with open chromatin and accessibility to transcriptional machinery\(^{79}\) linking butyrate to increases in gene expression and indirectly to decreases in DNA methylation\(^{75,80}\). Besides butyrate production, intestinal microbiota produce B vitamins and folate, metabolites that participate in one-carbon metabolism of the host, both as methyl donors (MD) and co-factors, potentially influencing the constitution of the methylome since they cannot be synthesised by humans\(^{81}\).

Corroborating the fact that the lack of bacteria in the intestinal tract can induce methylole modification of the intestinal epithelium, Yu et al. compared identical lines of mice in axenic or conventional (CV) conditions\(^{80}\). They observed an abnormal DNA methylation profile, specifically hypomethylation of CpG nucleotides in GF mice at 21 and 100 days of life, while on day 0, methylation patterns were indistinguishable between GF and CV mice. Interestingly, the hypomethylation was not global and did not affect genome repetitive elements, suggesting a specificity to the regulation of DNA methylation by the intestinal microbiota. Hypomethylated CGI in GF mice were related to lower expression of genes associated with intestinal maturation and that, for some genes, the expression level was increased to levels comparable to CV mice after the reconstitution of the communal bacteria by faecal microbiota transplant.

Pan et al. also investigated the influence of microbiota colonisation at different life stages on the methylome of small intestine epithelial cells\(^{49}\). They observed that the influence of microbial presence on DNA methylation was detected early after birth and global methylation level increased subtly over time. The number of differently methylated positions between CV and GF animals was ten times greater (1496 differently methylated positions) in the first week after birth than in week 4 and weeks 12/16, which corroborates with the idea that microbial colonisation has great impact on gut function during early life, when differently methylated positions were also enriched in promoter regions. However, adult mice (12/16 weeks) presented greater number of differently expressed genes containing differently methylated positions compared with only 17 in week 1. Furthermore, both Tet3 and Dnmt3a expression levels were altered by microbiota presence in week 1 and week 12/16 after birth.

Interestingly, the intestinal adaptation of preterm piglets to microbiota colonisation and milk enteral feeding was shown to involve DNA methylation changes and occurred mostly in the first month of life, with the global hypermethylation of preterm intestines observed within the first 5 days of life normalising to the levels of the term intestine by the 26th day after birth\(^{82}\). Methylation differences observed in the five postnatal days between preterm and term in mid intestine affected Wnt signalling and lipopolysaccharide-binding protein-toll-like receptor 4 (TLR4) pathways, indicating cell proliferation and immune alterations in the preterm gut.

In a more recent study, Ansari et al. observed a global hypomethylation in colonic crypts when comparing CV to GF mice accompanied by a subtle change in gene promoters\(^{83}\). Analysing low-methylated regions representing potentially active regulatory regions, they observed a greater number of
hypo- and hypermethylated regions (12985) in the CV mice with only 3115 hypermethylated when compared with the GF animals. Furthermore, the majority (78.2%) of low-methylated regions associated with changes in gene expression were hypo-methylated and from those, 300 were linked with significantly increased expression and were enriched for the binding sites of forkhead box A, kruppel like factor and activator protein 1 TCF. These findings suggest a much more intricate role of bacteria–host interaction on DNA methylation and regulation of gene expression. It also shows that the microbiota helps to regulate intestinal maturation and homeostasis through DNA methylation, possibly controlling TCF binding sites.

Takahashi and colleagues observed that the methylation profile of the TLR4 gene was different in CV and GF mice with the 5’ end of the gene being considerably hypomethylated in the intestinal epithelial cells of the large intestine of GF mice (84). Interestingly, Tlr4 expression was not associated with its methylation status in the small intestine, suggesting that the bacterial load in large intestine could be an important factor for bacterial regulation of Tlr4 expression and that there is a complementary regulatory process for Tlr4 expression in the small intestine.

Both microbiota shifts and difficulties in discriminating communal from pathogenic bacteria can play a role in regulating host functionality. Host recognition of bacterial populations and its link with DNA methylation, gene expression and host physiology were further corroborated with the use of TLR2 KO mice (23). The Tlr2 KO mice had different expression levels of genes associated with immune response correlated with changes in DNA methylation and were combined with shifts in colonic microbial populations with Firmicutes being less abundant and Proteobacteria and Bacteroidetes more abundant in the KO mice. Interestingly, the diversity was much higher for the bacteria present within the Tlr2 KO mice, compared with the much more closely clustered wild-type samples, indicating a more consistent control of microbiota population diversity by the presence of TLR2.

Interestingly, in a study from Bhat et al., neither probiotics nor E. coli exposure alone alters global DNA methylation in Caco-2 cells despite changes in histone acetylation (85). However, when cells treated with probiotic strains of Lactobacilli were challenged with E. coli, the result was global hypermethylation, indicating that interactions within microbiota populations also influence host functionality.

Although many studies focus on changes in microbiome composition due to different conditions, such as nutritional status (reviewed by Millon et al.; Castaner et al.; Blanton et al.) (86-88), diet (reviewed by Singh et al.) (89) antibiotic use (reviewed by Izumi et al.) (90) and pathological conditions (reviewed by Wang et al.) (91), only few look for associations between shifts in bacterial populations and DNA methylation status of the host despite observed changes in gene expression. Therefore, new research on the regulatory mechanisms, including DNA methylation, that bacterial populations use to impact host function can help to determine the important microbial–host interactions that influence physiological and pathological processes.

**Dietary supplementation and DNA methylation**

The association between diet and the intestinal microbial population has been extensively studied (92-95). Gene expression modifications and epigenetic alterations in different tissues and developmental stages due to host gene expression directly or indirectly associated with diet and nutritional status have also been extensively demonstrated (reviewed by Jiménez-Chillarón et al.) (94). However, the effects of dietary interventions on the intestinal tract epigenome are not well established and the molecular mechanisms regulating the change in transcription level in response to intake modifications or microbiota manipulations have still to be elucidated for a large part of the observed events. For instance, Krautkramer et al. demonstrated that the microbiota induces histone modification in the host tissue, including liver and colon, in a diet-dependent manner, suggesting modulation of bacterial population and function due to nutrient availability (94). This was confirmed by including SCFA to the diet of GF mice and observing a rescue of the expression level and chromatin modification to levels compared with colonised mice.

DNA methylation deposition requires not only the presence of DNMT enzymes but also the presence of methyl groups that are supplied by the one-carbon metabolic pathway (Fig. 2) that is dependent on nutrition (95,96). The major source of methyl groups for multiple cellular methylation processes (DNA, RNA, protein and lipid methylation) through the action of methyltransferases is S-adenosylmethionine, derived from one-carbon metabolism, which is dependent on multiple substrates, factors/co-factors such as B12, B6, folate, methionine and betaine (97,98). The intake of some of these nutrients, also known as MD, has been shown to have systemic effects on methylation, such as folate intake restoring methylation status of blood cells of patients with hyperhomocysteinaemia, a disease that characterised by an unbalanced one-carbon cycle that results in global hypomethylation (99).

Early nutritional effects on metabolism and disease predisposition have been observed on multiple occasions and extensively studied (94,109,101). One of the mechanisms by which nutrition can establish these long-lasting modifications is via DNA methylation as shown with the maternal MD supplementation of yellow agouti mice (102). The MD diet altered the coat colour of the offspring due to DNA methylation of the transposable element inserted in the agouti gene, indicating that dietary exposure in pregnancy or early life can have significant and long-lasting effects on metabolism and disease susceptibility in different organs of the offspring including the intestines. Policies for supplementation with folic acid for pregnant women to prevent open neural tube defects (103), the increase in the use of MD supplementation, especially in developed countries, and long-lasting effects of maternal diet on the DNA methylation of the offspring (104-109) encouraged the investigation into transgenerational risks and effects of maternal supplementation with MD (97).

Morphological changes due to MD supplementation/deficiency have also been observed in the intestines. Silva et al. found sporadic regions throughout the small intestine with increased crypt depth in the offspring of dams submitted to a
methyl deficient diet during gestation until weaning (110). The MD deficient offspring also had a decrease in jejunal and faecal microbial α-diversity and lower relative abundance of Bacteroidales and higher abundance of Lactobacillales. Interestingly, they also observed target hypomethylation of glycosylation genes in enteroids cultured in methyl deficient media (110). Furthermore, DNA methylation levels of two repetitive elements were measured and only one (IAP) was hypomethylated in the MD deficient enteroids, disregarding global hypomethylation and contradicting the one-carbon metabolism theory.

The one-carbon metabolism theory suggests that a reduced intake of dietary MD would induce a global hypomethylation due to decreased resources for formation of methyl groups, and the opposite would also be true, with an increase in DNA methylation due to a dietary supplementation of MD. This is corroborated by studies like the one from Choi et al., which describe a 35% reduction in DNA methylation in colonic genome of rats after vitamin B12 deficiency (111), and from McKay et al. that observed global hypomethylation in the small intestine of offspring of dams submitted to a low folate diet during pregnancy and lactation (112).

However, local hyper and hypomethylation were observed in studies with MD supplementation contradicting the idea that DNA methylation would decrease or increase in a random, dysregulated manner. Schaible et al., for instance, found that supplementation of the maternal diet with a combination of MD such as B12, folate and choline can alter the establishment of DNA methylation in mice as observed at postnatal day 30 (P30) and P90 (24). They observed fifty-nine hypermethylated regions and ninety-six hypomethylated ones, with more than 50% hypomethylated loci localised on the X chromosome. The changes in methylation were combined with an increase in the susceptibility to acute colitis induced by dextran sodium sulphate associated with changes in gene expression. Eighteen of the 155 DMR were associated with change in expression of a flanking gene, including the PPARα gene, which showed decreased methylation level and increased expression. Finally, they also assessed the mucosa-associated microbiome of the offspring and found a significant separation at the genus level at P30 that is rescued by P90 with Bacteroidetes and Clostridia being overrepresented, and Lactobacillus underrepresented in early life of MD-exposed offspring.

Maternal diet can also influence the DNA methylation level of the PPARα promoter in the liver of the offspring (113,114), with protein-restricted diet having 26% less methylation than the controls and folic acid rescuing the methylation status (114), corroborating with the findings of Schaible et al. In the murine fetal gut, a low folate maternal diet reduced the methylation in the Zn transporter gene solute carrier 39 member 4 (Slc39a4), but did not cause changes in the methylation of insulin-like growth factor 2 and oestrogen receptor 1 (115). Paternal folic acid deficient and supplemented diet throughout life also impacted DNA methylation levels of imprinted genes in the brain and in the placenta and increased postnatal–preweaning pup death (116), as was observed by Ly et al.

Despite the fact the important epigenetic modification occurs in colonic mucosa during paediatric development as shown by Kellermayer et al. (22), Schaible et al. did not observe an increase to colitis susceptibility when the mice were exposed to MD supplementation from P30 to P80, suggesting that the in utero effect is more important than postnatal dietary exposure. These results were in agreement with the findings from Mir et al. that, in spite of not analysing the DNA methylation status, observed a 25% increase in mortality in the MD group offspring, with males being more susceptible to colitis (117). The sex difference might be
related to the finding of Schaible et al. that a high percentage of hypomethylated regions due to maternal MD were localised on the X chromosome. Trasher and colleagues also did not observe changes in global methylation in the colon of mice predisposed to intestinal adenoma formation (Apc^{−/+Min}) submitted to a low fat diet from weaning to 13 weeks of age with or without reductions in Dnmt1 expression due to heterozygous gene KO[119].

It is noteworthy that the impact of methyl deficient/supplemented diets on DNA methylation is organ specific, as observed in studies focusing on liver and brain DNA methylation in rats under dietary restriction of methionine, choline and folic acid. In these studies, global hypomethylation was observed in the liver[119] and genome hypermethylation in the brain[120]. Ly et al. 2016 also observed that maternal supplementation with folic acid during the second and third week of gestation or throughout pregnancy decreased global methylation in the brain, while no global alterations were observed in the liver, colon or the kidney of the offspring pups[121].

The tumour suppressor gene p53, generally mutated or dysregulated in cancer, showed an increase in DNA methylation in its promoter in liver and colon mucosa following selenomethionine diet supplementation[122]. Global DNA methylation was decreased in the liver of rats fed with extra selenomethionine, while the selenomethionine-deficient animals had global hypermethylation. This was not true for colon mucosa, where the selenomethionine intake did not significantly influence global methylation status. Using a modelling approach to identify variables that had the strongest association with gene-specific methylation, Tapp et al. observed greater correspondence between age and methylation level in males than in females and that correlation was weak and slightly negative for LINE-1 elements, indicating a subtle age-related hypomethylation[123]. Plasma folate and red cell folate correlated with the methylation of some of the genes in a sex-dependent manner. A similar trend was observed with plasma selenium that presented a positive correlation with methylation in males but a negative correlation in females. Interestingly, neither B12 nor homocysteine had a significant correlation with methylation status.

Other diet supplementations, not directly associated with one-carbon metabolism, have also been shown to influence DNA methylation in intestinal tissue. One example was the association of epigallocatechin-3-gallate, the catechin of green tea, to a regulatory effect on DNMT enzymes[124]. It has also been shown to inhibit DNMT in multiple cells including colon cell lines[125]. Despite that the majority of epigallocatechin-3-gallate is absorbed in the small intestine, the administration of epigallocatechin-3-gallate resulted in a shift in microbiota population in the colon of mice, resulting in a decrease in acetic and butyric acids[126] suggesting a modulation of energy metabolism. Remely et al. found that the methylation of CGI associated with the promoter region of mult1 homolog 1 (MLH1) in the colon increased with ECGC intervention to a greater extent than with a high fat diet, with or without ECGC[127]. In fact, a high fat diet coupled with ECGC decreased the methylation level of this CGI. Dmnt1 gene expression was lower in the colon of mice fed a high fat diet, but the expression level was rescued when ECGC was added to the diet. CpG in the promoter region of Dmnt1 were hypermethylated when ECGC was added to either high fat or control diets.

Organ-specific effects were also observed with dietary supplementation of nutrients outside one-carbon metabolism. Day et al. observed DNA methylation changes in the prostate of rats but not in the liver after diet supplementation with the soy phyto-oestrogen genistein[128]. This was similar to the findings of Guerrero-Bosagna et al., where continuous pre- and postnatal genistein and daidzein exposure did not alter hepatic methylation level in the promotor region of the gene a-actin (Acta1), but it caused hypermethylation in the pancreas[129].

All this evidence suggests that the diet can affect the methylation status systemically and locally in the intestinal tract, representing both a risk and a protective function for specific disease pathologies. The consequences of food and nutrient intake on the molecular mechanisms regulating gene expression and consequently the function of the intestines should be further studied as an easy and affordable way of preventing and treating pathological conditions/states.

**Nutritional status and DNA methylation**

Extreme nutritional status, such as undernutrition and obesity, has been reported to influence DNA methylation in different organs with a potential impact on the health of individuals as parental dietary conditions also effect the metabolism and methylation level in offspring. Maternal nutrition has been shown to influence the state of DNA methylation in offspring in several animal models. In rats and mice, it has been reported that maternal malnutrition due to protein deficiency resulted in stable global hypermethylation until adulthood in the liver of the offspring[129]. Methylation changes at specific loci in the liver[113,114,130–138] pancreas[139], amygdala[140], adrenal gland[141,142], hypothalamus[143] and adipose tissue[144,145] were also reported correlating with changes in gene expression. Likewise, low protein feeding caused paternal transgenerational effects in mice, including various DNA methylation changes in the offspring’s liver[137]. In utero undernutrition increased obesity and glucose resistance in mice and interestingly, the female offspring of male mice submitted to in utero undernutrition also developed glucose intolerance which was linked to DNA methylation alteration on the liver X receptor alpha (Lxra) gene in the liver, the same methylation signature found on the sperm of the F1 mice[135]. Accordingly, Radford et al. found that in utero undernutrition changes the germine methylome of the male offspring with a prevalence of hypomethylation and enrichment of DMR in nucleosome retaining regions, and part of the alteration is resistant to embryo methylation reprogramming[140]. However, Ivanova et al. found that the liver DMR identified in mice treated with maternal or early postnatal protein restriction did not significantly affect imprinted genes, indicating that expression of imprinted genes is not particularly influenced by maternal or early life nutrition[147].

Maternal diet during pregnancy was also shown to impact methylation status in species other than rodents. The offspring of sows under dietary protein restriction during pregnancy developed changes in DNA methylation in CGI flanking
metabolic genes, such as PPARα, in the liver\(^{148}\). Maternal low-protein diet also affected mitochondrial DNA methylation in the liver of piglets\(^{149}\) and muscle GLUT type 4 (GLUT4) promoter methylation level\(^{150}\). Energetic restriction in baboons during early pregnancy resulted in global hypomethylation in fetal kidneys and in late gestation resulted in global hypermethylation in fetal kidneys and frontal cortex\(^{151}\). Hepatic epigenetic modification was also observed in baboon fetus due to maternal malnutrition\(^{152-154}\). Tobi \textit{et al.} found hypomethylation at the insulin-like growth factor 2 locus and other genes in leukocytes of women that were affected by famine during the peri-conception period and that the difference of methylation in some of the genes was sex specific\(^{155-157}\).

Dietary interventions can also influence epigenome changes in early/adult life specially during dietary transitions. Post-weaning malnourishment in mice caused global hypomethylation in the thalamus and hypothalamus with methylation profiling in the thalamus identifying DMR (both hypomethylated and hypermethylated) in genes associated with neuronal development or psychiatric diseases, including nine genes related to long-term potentiation\(^{158}\). Peter \textit{et al.} found associations between the blood DNA methylation profile associated with severe malnutrition in early childhood and impairments in attention and cognition\(^{159}\). A study in India searched for correlation results between DNA methylation in LINE1 sites in DNA from blood cells of children from 5 to 12 years old with malnourishment indices such as Z-score, BMI and blood vitamin concentration (B12 and folic acid) and found that the DNA methylation at LINE1 elements relates inversely to retinol levels in the blood\(^{160}\).

Ageing is usually associated with global hypomethylation combined with hypermethylation at specific loci, which is also the methylation profile linked to cancer (reviewed by Klutstein \textit{et al.})\(^{161}\). Energetic restriction can potentially reverse age-related abnormal DNA methylation, increasing genomic stability\(^{162,163}\) in a possible explanation as to why it is considered the most powerful mechanism to increase lifespan in different animal models\(^{162,164-166}\). Rhesus monkeys exposed to chronic 30% energetic restriction had a reduction in the age-related methylation drift in their blood compared with \textit{ad libitum} fed controls\(^{167}\). The same study found that in mice with 40% energetic restriction, similar but more evident results were observed across multiple tissues. Furthermore, fifteen genes that showed an age-related drift in methylation in the blood of the monkeys were analysed in various tissues including the small and large intestine, and an even larger methylation drift was seen in the large intestine with the selected genes also showing age-related clustering in both the small and large intestines indicating a marked effect of diet on the DNA methylation profile in these tissues.

High-fat diets and maternal overnutrition and obesity during gestation also influence the DNA methylation levels of the offspring in tissues including the brain\(^{170,171}\), liver\(^{172-174}\), blood\(^{175}\) and adipose tissue\(^{176}\). Contrastingly, Li \textit{et al.} found that offspring of obese female mice presented widespread but subtle alterations in their hepatic DNA methylation profile and suggested that healthy postnatal feeding would be enough to prevent metabolic dysfunctions in the offspring\(^{177}\). This is in agreement with Moody \textit{et al.} that showed evidence that postnatal diet can reverse the methylation effects in the liver caused by a high fat maternal diet\(^{178}\). As with undernutrition, in rats, female offspring of high fat-fed males presented with programmed β-cell dysfunction despite the fact of not being exposed to a high-fat diet during development\(^{179}\) and the alteration was accompanied by DNA methylation changes. There is also evidence that obesogenic diets combined with obesity-associated microbiome modulate colonic gene expression through epigenetic modifications other than DNA methylation\(^{180}\). Maternal and paternal obesity have also been associated with DNA methylation changes in fetal blood and placenta at specific loci\(^{181-183}\).

Chronic high-fat diet in mice (from weaning to 20 weeks of age) changed promoter methylation status of genes associated with food intake in brain tissue\(^{178,184}\), which could explain the establishment of obesity and obesity-related diseases. Neonatal overfeeding has also been associated with changes in DNA methylation in the hypothalamus of rats and was associated with decreased expression of genes associated with the metabolic syndrome\(^{171,184,185}\). There has also been work relating DNA methylation profiles (especially in blood) with BMI, waist circumference and body composition. In an epigenome-wide study using blood samples from human subjects\(^{186}\), it was observed that validated, differently methylated CpG associated with genes or coding regions explained 14.18% of the BMI score and 16.73% of the waist circumference variability. Interestingly, from the 95 loci with CpG significantly associated with BMI, only ten were genes previously associated with BMI from a genome-wide association study. Using blood samples from preschool children, Rzehak \textit{et al.} found specific DNA methylation variants associated with BMI, fat mass and fat-free mass\(^{187}\).

Noticeably, the link between extreme nutritional status and developmental programming has been extensively studied, especially in the liver, brain and adipose tissue, while information regarding the intestines is still lacking. However, if nutritional supplementation and microbiome colonisation have organ-specific effects, it is not farfetched that the effects of obesity and undernutrition in these tissues do not reflect what would be observed in the intestinal tract and the different sections of the small and large intestine.

Although morphophysiological changes in the intestines due to malnutrition have been shown\(^{188}\), including in swine models for childhood undernutrition\(^{189}\), the molecular aspects underlying these changes have not been understood. Furthermore, there have been few efforts towards analysing DNA methylation profile changes in the intestines, especially the small intestines, due to extreme nutritional status. As the point of adaptation between nutritional intake and immune protection, changes in the intestinal tract could potentially be related to systemic effects of dietary manipulations. The known effects of nutritional status on the epigenome of the intestinal tract are limited and could be the key to explain how dietary intervention can systemically influence host physiology.
Conclusions

It is easy to see how information on the molecular effects of how diet and nutritional status directly influence the intestine and how the interplay between nutrient intake/nutritional status and commensal bacteria influences host health and their long-lasting effects on metabolism and disease predisposition is still lacking. The use of diet to manipulate host physiology to better respond to pathological or adverse conditions such as endo-mergenon, chronic inflammatory diseases and diarrhoea could be used once the mechanisms behind the interactions between diet, microbiota and host are understood. As observed in this review, DNA methylation can help to link environmental shifts to host function, but there is still a long way to go in describing the actions of bioactive nutrients and foods that could benefit host health. The links between intestinal and nutritional biology need to be better defined in order to answer clinically relevant questions. For instance, could diet/microbiota modulation be an effective and precise approach to manipulate intestinal cell function against specific medical conditions? At what stage of life would these alterations need to occur and are they sustainable over time? Specifically, the role of epigenetics as a conductor makes it a crucial point of study to deepen our knowledge of intestinal adaptation to nutritional challenges and discover its translational potential in precession medicine.

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