Tasks of some dietary essential nutrients on epigenetics and one carbon metabolism in livestock during production

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Abstract: Interchanges in the epigenome caused by the environment have been cited in multiple animal being, ranging from insects to rodents passing through poultry, larger animals, and nonhuman primates to humans. These include DNA methylation, chromatin remodeling, histone modifications, and more recently the index has magnified to incorporate noncoding RNAs and microRNA gene regulation. Thus, it is increasingly accepted that, nutrition one of the factors that alter gene expression and affect's health during animal productivity through epigenetic modification. DNA methylation is the most widely studied form of epigenetic modification and occurs within the one-carbon metabolism tract which is dependent upon a number of enzymes in the presence of micronutrients, including folate, choline, and betaine acquired through the diet. In vertebrates DNA methylation is primarily a stable repressive mark built at cytosines in CpG dinucleotides; however, its regulation is more powerful than previously believed. Nutri-Epigenomic is an emerging discipline examining the role of nutrient on gene expression. Ultimately, DNA methylation and other epigenetic status, as well as dietetic practices, particularly micronutrient intake, may influence health and productive phenotypes, also playing as significant biomarker in additive-benefit assessment during the period of use. There are a tight interrelationship betaine, choline and methionine. All are dietary sources of methyl groups made it prominent when studying diet-DNA methylation. In this review, our team will focus on the role of important methyl donors and nutrient-gene inter actions through the one-carbon metabolism.

Keywords: Nutri-Epigenomics, Livestock, methyl donors, one carbon metabolism.

I. Introduction

Since 1940s the term epigenetics was first proposed by Waddington, where, it was early used to describe a new phenotype due to gene-environment interaction. That way indicative the source of the phenotype from the genotype, a procedure that is also affected by the environment. As from 20 years ago, epigenetic trends had become in increasing way every day, sharing new sciences serving human health, pathogenesis of diseases, embryogenesis, agriculture sciences, including plant and animal productivity as well as animal models used to enhance the human researches. Many definition regarding epigenetic, were discussed in many articles with the same strategy of changes in gene function (expression), with new phenotype due to changes in gene physiology other than anatomical (sequence) feature. Epigenetic, is investigating and analyzing mitotically and/or meiotically inherited changes in gene function, that are away from changes in gene sequence\textsuperscript{1,2}. Also, “Any long-term change in gene function that does not involve a change in gene sequence or structure”\textsuperscript{3}. There are a continuous conversation among cellular elements and the surrounding environment, epigenetic modifications are dynamic throughout the lifecourse and can be heavily influenced by external factors, and thus, external effects on the epigenome may alter gene expression, potentially giving rise to new phenotype. In a brief, DNA and chromatin alterations that remain from a cell division to the subsequent one, regardless of changes in the basic DNA sequence, known as epigenetics. Whereas, the in general epigenetic status of a cell that suits as an interface between the genome and the surround environment referred as epigenome. The dynamic character of epigenetic rule, unlike the static nature of the gene sequence allows a mechanism for reprogramming gene function in response to environmental changes, including diet. Thus, epigenetics might give an interpretation for well recorded gene x environment interactions\textsuperscript{3}. Diet and its fine elements are the most environmental factors controlling intensity of gene expression, for example, honeybees grow to be either queens or workers relying on whether they are fed royal jelly or bee bread. Despite being genetically equal at the larval stage, honeybees fed pure royal jelly are obviously different from worker honeybees. The different honeybee phenotype occurs as a result of epigenetic modifications in DNA methylation styles induced by the kind of honey they are fed with, therefore, showing a strong link between diet-induced epigenetic modifications that lead to unusual development from the same genetic background. In addition, other reports in mammals revealed that environmental stimuli - different dietary factors- can modify DNA methylation and histone acetylation models and, accordingly, influence gene regulation and the phenotypical expression of gene\textsuperscript{4}. Some of the dietary micro-elements were revealed documented phenotypes due to their epigenetic effect, so in this review we would show basic epigenetic machineries beside the essential role of some methyl donators that affecting gene expression away from changes in gene sequence.

II. Epigenetic machineries

DNA as well as non-coding RNA methylation and histone post-translational modifications are important epigenetic mechanisms regulating gene activity.
Role of some dietary methyl donors affecting epigenetics and one carbon metabolism during ....

2.1 DNA Methylation
In mammals, methylation of DNA is an epigenetic adaptation that commonly exists at 5-methylcytosine within CpG (C followed by G) dinucleotide (Regions of the genome rich in CpG dinucleotide often located at the promoters of genes) and it is an essential tool for epigenetic regulation of the genes expression. Methylation of CpG within these regions is generally associated with gene repression [5, 6]. Oftentimes, CpG methylation within the gene body itself and outside of CpG islands of gene resulted in its activation [7]. DNA methylation patterns are essential for mammalian development and are faithfully maintained through cell division by DNA methyltransferases [8]. This DNA methylation can silence the gene expression and is imperative in keeping genome stability. Increasing risk for a disease onset was recorded when these epigenetic patterns are not maintained [9]. Methytransferases and demethylases are family of proteins (enzymes) that add or remove methyl group to the cytosine of the newly synthesized DNA. Four methyltransferases were identified in human and mouse, DNAmethyltransferase3 (DNMT3) family Dnmt3a, and Dnmt3b accountable for setting up methylation of DNA and DNMT1 keeping up the DNA methylation, but DNMT2 was found to have little methyltransferase activity, by in vitro trials, all the DNA methyltransferase mRNA-sequences cloned in bovines, sheep and chicken [10]participate very strong homology with those of mouse and human, that shows a conservative role of DNA methyltransferase in various animals. Different isoforms of Dnm1t1, Dnmt3a, and Dnmt3b were also found in bovine; even so the functions of these isoforms in DNA methylation stay unknown [11, 12].

2.2 Histone modifications
The network of DNA and histone proteins that can be built within the nucleus of eukaryotic cells define the chromatin. It specifies a scaffold for the wrapping of the whole genome. Nucleosome is the primary working unit of chromatin; it contains 147 base pairs of DNA, which are wrapped all over a histone octamer that exists in two copies each of histones H2A, H2B, H3 and H4. Unusual DNA, the histones can be covalently adapted in different manners including acetylation, methylation, sumoylation, phosphorylation, and ubiquitination [13].

2.3 Chromatin remodeling
The gene expression is influenced essentially by chromatin structure. The euchromatin and heterochromatin in the eukaryotic genome designate for the transcriptionally active and inactive domains, separately [14, 15]. DNA methylation [16], histone modification[17], and ATP-dependent chromatin-remodeling structure [18] are three major factors that govern the assemblage and normal of chromatin structure.

2.4. RNA epigenetics
Cellular RNAs carry diverse chemical modifications that used to be regarded as static and having minor roles in ‘fine-tuning’ structural and functional properties of RNAs [19]. The recent development of novel transcriptome-wide approaches to capture global m5C and m6A RNA methylomes has not only restored scientific interest in the field but also contributed to a better understanding how gene expression is regulated at different levels. The importance of a tightly controlled deposition of both m5C and m6A into RNA is further underscored by the strong link of loss-of-function mutations in methylating and demethylating enzymes to several severe human diseases [20]. Recent studies have discovered protein ‘writers’, ‘erasers’ and ‘readers’ of this RNA chemical mark, as well as its dynamic deposition on mRNA and other types of nuclear RNA. These findings strongly indicate dynamic regulatory roles that are analogous to the well-known reversible epigenetic modifications of DNA and histone proteins[21].

III. Methyl donors and methylation processing
Methylation of CpG dinucleotide orders methyl donors from dietary elements suchas folate, choline and betaine, which play key roles in restoring methionine (MET) following transmethylation, the transfer of a methyl (CH3) group from the stimuluted form of MET (S-adenosylmethionine; SAM) to a donee molecule. S-Adenosyl-methionine is synthesized from methionine by S-adenosylhomocysteine synthase (also known as methionine-adenosyl-transferase [22]. Transmethylation is needed in at least 100 various metabolic reactions [23] and is necessary for various metabolic actions including regulation of DNA expression, muscle contraction, hormonal signaling, neurotransmission, cell membrane integrity, cell growth and protein synthesis. It is notable that the multiplication of lymphocytes and macrophages is also influenced by methylation reactions. The product of transmethylation, S-adenosylhomocysteine (SAH), is changed to homocysteine (Hcy), that can either be irreversibly lost from the cycle by trans-sulfuration to cysteine (CYS) or remethylated to MET by using methyl groups obtained from 5-methyl-tetrahydrofolate (5m THF), a byproduct of folate, choline or betaine. Under normal situation, diminished store of MET to the cycle will decline SAM concentration which will favor remethylation by reducing SAM-mediated inhibition of Methylenetetrahydrofolate reductase (MTHFR) and Betaine-homocysteine methyltransferase (BHMT), and decrease trans-sulfuration by withdrawing Cystathionine beta synthase (CBS) activation. A lack of methyl donors will have the same impact on these enzymes, and HCY will aggregate because of a lack of sufficient methyl groups for remethylation and low CBS activity [24].

Some essential roles of some methyl donors in animal body.
The first methyl donor for DNA methylation is S-adenosylmethionine (SAM), a type generated in the cyclical cellular procedure called one-carbon metabolism (transfer). One-carbon metabolism is catalyzed by several enzymes in the presence of dietary micro nutrients, including folate, choline, betaine and other B vitamins. For this reason, nutrition status, particularly micronutrient intake, has been a focal spot when studying epigenetic mechanisms. [25].

3.1. Methionine (Met)
From the nutritional point of view, Met is classified as essential (carbon skeleton cannot be synthesized by the body) sulfur-containing AA that tasks many roles in the body including(1) participation in protein synthesis and the

DOI: 10.9790/2380-0908028690 www.iosrjournals.org 87 | Page
production of other sulfur-containing amino acids (e.g. homocysteine – a sulfur-containing AA which is an indirect product of methylation and trans-sulfuration, in what is called recently “protein related role of AA” [26, 27], (2) acting as a precursor of carnitine and glutathione, thus protect cells against oxidative stress [28, 29, 3] labile methyl group (CH3) is incriminated in the metabolism of energy, which have to be provided from the diet [30]. Under the influence of methionine-adenosyltransferase, Met is converted to S-adenosyl-methionine (SAM). S-adenosyl-methionine is a common co-substrate that supplies methyl groups required for various metabolic processes, including DNA methylation and synthesis of RNA, proteins and lipids. In the process of methylation, SAM is converted to S-adenosyl-homocysteine in the presence of methyltransferases “non-protein related role of AA” [32, 31]. Cysteine (Cys) is grouped as semi essential because it can be produced from Met [32]. The concept of alternate dietary methyl donors (i.e. betaine, choline, methylneogenesis precursors) sparring the methionine requirement has recently been discussed [33]. The inverse association between plasma betaine and homocysteine levels was most noticeable when serum folate levels were low [34]. Because -in human- betaine [35], choline [36] and folic acid [37], be able effectively y lessen plasma homocysteine by enlarging remethylation to methionine, these nutrients are very likely to be able sparring the methionine demand. Actually, in smart- isotope studies the choline and folic acid doubled remethylation to methionine in methionine deficient diets. That idea can be applied by using fewer costly additives containing choline and/or betaine in poultry diets. Several studies proved that betaine and/or choline can reserve methionine demand in growing chickens [38] and young pigs [39], not only for the synthesis of protein but also for creatine and carnitine synthesis. The fact that these studies were able to show a sparing effect of methyl groups on the methionine requirement demonstrates the enormous demands of these pathways.

3.2. Choline and betaine (N,N,N-trimethyl-glycine)

Choline and betaine are dietary leading nutrients with several biological functions. Along with folate and other B vitamins, choline and betaine contribute to one-carbon metabolism that constitutes a network of integrated biochemical pathways transferring methyl groups from one compound to another [40, 41]. Choline is oxidized to betaine, both folate and betaine donate methyl groups to homocysteine and produce methionine, which transfer a methyl-group to S-adenosyl-methionine (a universal methyl-group donor) for methylation of DNA and RNA. Through this process, one-carbon metabolism affects genomic stability and expression and intervenes nucleotide synthesis. Thus, a number of animals in vivo and, to a lesser extent, human studies have investigated the role of dietary choline and/or betaine and their impact on global and candidate gene DNA methylation. As, choline-deficient diet lessens hepatic folate levels in animals [42]. Parallel, a folate deficient diet lowered hepatic choline levels in animals [43]. A double-blind clinical study of folate supplementation raised plasma betaine levels in a dose-dependent style following folate supplement [44]. Plasma choline levels existed reactive to reduction and subsequent replention of folate intake [45]. Betaine and choline have additional biological galas not shared by each other. Choline is a precursor for phosphatidylcholine and sphingomyelin, which are integrated into cellular membrane and involved in signal transduction [46]. Choline is also a pioneer for the neurotransmitter acetylcholine and is concerned in brain development and normal memory job [47, 48].

Betaine is an osmolyte (play a role in maintaining cell volume, fluid balance and protects cells, proteins, and enzymes from water stress[49, 50]. Betaine supplementation plays crucial roles in nutritional programming through epigenetic regulation of gene expression in pigs, ducks meat and egg type chickens. Betaine supplement was associated with retained positional effects in the -rate limiting enzyme- lipoprotein lipase (LPL) gene promotor, as measured in adipocytes [51]. With addition of betaine, most of the promoters were highly hypo-methylated and many CpG sites were built to be hyper methylated. Interestingly, LPL gene expression was significantly decreased with betaine supplementation. Maternal betaine supplementation enhances betaine/methionine metabolism and DNA methyltransferase expression, causes hypermethylation of the differentially methylated region (DMR) on IGF2 gene, which was associated with augmented expression of insulin like growth factor II (IGF2) and cell proliferation/anti-apoptotic markers in the hippocampus of neonatal piglets [52]. Dietary betaine supplementation (3g/kg diet) during gestation of sows, attenuates hepatic lipogenesis in neonatal piglets via epigenetic and GR-mediated mechanisms [53]. Since one-carbon (1C) metabolism consists of an integrated series of metabolic pathways that include the folate cycle and methionine remethylation and trans-sulfuration pathways, maternal nutrient status can cause epigenetic alterations to the genome of the developing fetus, which potentially can impact future generations [54].

3.3 Folic acid (Folate)

folate (B9), riboflavin (B2) cobalamin (B12), serve as cofactors in (1C) metabolism while pyridoxine (B6) is essential for transsulfuration pathway, folate play as a catalytic substrate for the transfer of (1C) units [25, 32]. Numerous reports have indicated a link among folate, choline, and lipid metabolism. Folate has a role in the maintenance of cellular S-Adenosyl-methionine (AdoMet) and Adenosyl-homocysteine (AdoHcy) levels. Dietary folate play a role in epigenetic mechanisms, in the liver, that responsible for many methylation reactions which are used for post translational modification of proteins, methylation of DNA, and the synthesis of hormones, creatine, carnitine, and phosphatidylcholine PC. The tetra-hydro-folate (THF) plays a crucial role in a number of reactions that generate methyl groups from the catabolism of sarcosine, serine, dimethylglycine, and glycine [6]. Recently, folate status may be influencing microRNA expression linked to the severity of fatty liver disease. MicroRNAs (miRNAs) are short, non-coding transcripts, of closely 22 nucleotides in length. They belong to a regulatory class of RNAs which repress expression of target mRNA. Folic acid supply controls the expression of miRNAs perhaps by changes in methylation levels of promoter regions in the genome [55]. The severity of Non-alcoholic fatty liver disease (NAFLD) induced by a choline- and folate-deficient diet in mice is associated with altered expression of hepatic miRNAs, including miR-181a, miR-34a, miR-200b, and miR-221 [56]. Moreover, maintenance and regulation of the epigenetic state which depends on (1C) metabolism require adequate provision of B vitamins (including folate, vitamin B6 and vitamin B12), glycine, methionine, serine, choline, histidine, and creatine [57]. These nutrients play an important role in regulating the availability of S-adenosyl-methionine (a major methyl donor for DNA and protein methylation by specific
Role of some dietary methyl donors affecting epigenetics and one carbon metabolism during ....

DNA and protein methyltransferases [58]. For example, essential B vitamins, folate, and methionine deficient diet during the breeding period in sheep resulted in altered DNA methylation, insulin resistance, and higher blood pressure mostly observed in adult male kids [59].

3.4. Vitamins B2, B6 and B12
The water-soluble vitamins B2, B6 and B12 have an essential catalytic role in folate and one-carbon metabolism. Vitamin B6 serves as a coenzyme to serine hydroxyl-methyltransferase, the vital enzyme in the folate cycle converting THF to 5,10-

methylenetetrahydrofolate [60]. Riboflavin, or vitamin B2, is a precursor for Flavin-adenine-dinucleotides, which is a cofactor to MTHFR, the enzyme responsible for the reduction of 5,10-methylene THF to 5-methyl THF[61]. Vitamin B12 is the coenzyme of methionine synthase, which catalyzes the reaction of homocysteine, the by-product of SAM, to methionine. Therefore consumption of these water-soluble B vitamins has the potential to affect the efficiency of the one-carbon metabolism pathway. Both animal and human studies have evaluated the role of vitamin B12 in DNA methylation profiles. An in vivo rat model showed that deprivation of vitamin B12 in addition to a diet supplemented with folate enhanced placental global hypo-methylation compared to a diet exclusively supplemented with folate, suggesting that the interaction between micronutrients can alter methylation patterns more profoundly than excess or deprivation of just one micronutrient [62]. vitaminB12 and folate supplemented rat maternal diet, increases methylation of DNA of the agouti gene in the offspring, preventing the development of obesity [63]. The effect of maternal diet supplementation with methyl donors can be succeeded to the next era through germline epigenetic modifications [64]. Consequently, the nutritional and physiological state of the animal will alter the production of these bioactive substances, thus regulating the availability of methyl donors. When cystine or taurine is deficient in the diet, their synthesis from methionine will be increased in vivo, thus decreasing total S-
adenosylmethionine availability for DNA or protein methylation. Inadequate synthesis of glycine and serine, coupled with low supplies from the diet, can also impair (1C) metabolism [65]. So, an essential AA deficiency can alter the epigenetic state by changes in DNA methylation as well as histone modifications [66].

IV. Closing Remarks
Biochemical interactions can direct Cross-talk between nutrients (micronutrients) and gene expression. Because methylation is a cell type dependent, a comprehensive epigenetic analysis remains an optimal approach. Future studies should continue to focus on tissue specificity in DNA methylation investigation. To apply the rational and economic use of the most expensive micro-nutrients in the animals and poultry diets, adoption of studying nutri-epigenetics in production trials might help understanding productive phenotypes and re-estimation of dietary requirements of essential nutrients. One-carbon metabolism role perform an outstanding role in DNA synthesis and in validating and maintaining the epigenetic sketch via their diverse metabolic intermediates and contributing IC units. Thus, an impaired 1C metabolic pathway may cause changes to intracellular methyl pools that limit the availability of methyl groups for relevant DNA and histone modifications. There are two interacting factors that can affect this methyl paddle 1) the dietary provision of methyl donors (folate) and 1C metabolic cofactors (vitaminB12), 2) nature and extent of polymorphic variations in genes encoding 1C metabolic enzymes. Abnormal 1C metabolism can lead to aberrant methylation patterns in DNA and associated histone proteins that affect genome stability, gene expression, cellular differentiation and long-term development [67]. Either under nutrition or over nutrition during pregnancy (peri-conceptional time), remains a important problem in both medicine and animal agriculture as it can results in epigenetic modifications of some genes in both animals and humans. These changes are influenced by various factors (severity of malnutrition, sex, and gestational period) and may persist in offspring during postnatal lifetime and may transport to the next generation. Nutrients, particularly amino acids and B vitamins, are essential for the regulation of epigenetics. Binding evidence indicates that, the fetal and early neonatal ages of development are radically sensitive to environmental implies that have long-lasting consequences’ to postnatal growth, health, and likely athletic performance.

Conflict of interest statement
The authors declare that there are no conflicts of interest.

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Role of some dietary methyl donors affecting epigenetics and one carbon metabolism during ...


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