Alternative pathways for hydrogen sink originated from the ruminal fermentation of carbohydrates: Which microorganisms are involved in lowering methane emission?

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Abstract
Agriculture is responsible for a great share of the anthropogenic sources of greenhouse gases that, by warming the earth, threaten its biodiversity. Among greenhouse gas emissions, enteric CH_{4} from livestock is an important target to slow down climate changes. The CH_{4} is originated from rumen fermentation and its concentration is affected by several factors, including genetics and nutrition. Ruminants have an extraordinary symbiosis with microorganisms (bacteria, fungi, and protozoa) that ferment otherwise indigestible carbohydrates, from which they obtain energy to grow and continue actively producing, among other products, volatile fatty acids, CO_{2} and H_{2}. Detrimental ruminal accumulation of H_{2} is avoided by methanogenesis carried out by Archaea methanogens. Importantly, methanogenesis is not the only H_{2} sink pathway. In fact, other bacteria can reduce substrates using metabolic hydrogen formed during carbohydrate fermentation, namely propionate production and reductive acetogenesis, thus lowering the CH_{4} produced. Although the complexity of rumen poses challenges to mitigate CH_{4} production, the emergence of sequencing techniques that allow the study of microbial communities, gene expression, and metabolome are largely contributing to unravel pathways and key players in the rumen. Indeed, it is now recognized that in vivo emissions of CH_{4} are correlated to microbial communities, and particularly with the abundance of methanogens, several bacterial groups, and their genes. The goal of CH_{4} mitigation is to work in favor of the natural processes, without compromising rumen function, animal health, and productivity. Notwithstanding, the major challenge continues to be the feasibility and affordability of the proposed solutions.

Keywords: Acetogenesis, H_{2} sink, Methanogenesis, Microorganisms, Propionate, Rumen fermentation

Introduction
Global warming threatens biodiversity alongside the life of humans. Among other gases, the release of CO_{2} and CH_{4} into the atmosphere contributes significantly to the greenhouse effect, a phenomenon that prevents the reflection of solar energy back from the earth’s surface, causing a rise in temperature [1]. Agriculture is responsible for a great share of the anthropogenic sources of greenhouse gases (GHG) [1]. According to FAO (2013), GHG emission from livestock represents 14% of human-induced emissions, being beef and dairy cattle the main contributors [2]. Within the livestock sector, feed production, processing, and transportation account for \approx 45% of total GHG, followed by enteric CH_{4} emissions (\approx 40%) [2]. Therefore, acting over this latter source constitutes an opportunity to achieve the goals of the Green
Deal, which seeks for a 55% cut in GHG emission by 2030, compared to 1990 levels [3]. Although animal farming is often cited as the cause of the problem, it can actually be part of the solution. If other anthropogenic activities (e.g., burning of fossil fuels) slow down the emission of CO₂, mitigation of GHG emission, or at least the share from the livestock sector, might be achieved by sequestration of carbon in grasslands [4]. Carbon from the atmosphere (CO₂) is fixed in the soil mainly through plant photosynthesis and thus converted into organic material (e.g., grasses and forages), which is then consumed by grazing animals [5]. From the ruminal fermentation of carbohydrates, CH₄ is produced at the expense of energy and released into the atmosphere, where after approximately 10 years, it is broken down and converted back into CO₂, giving continuity to the natural carbon cycling, in a process called the biogenic carbon cycle (Fig. 1) [6]. Therefore, in theory, the reduction of animal production, which entails a decrease in protein availability, is not the only alternative to tackle GHG emission. Instead, enhancing animal productivity while decreasing CH₄ might be a sustainable option, not compromising feed for a growing population. In the latest years, a lot of effort has been put into the study of animal breeding [7], vaccines [8], dietary management, and additives [9], as means to mitigate CH₄ emissions. Despite positive results reported in some studies, there is still not a consensus at a global scale, mainly because the efficacy and feasibility of each proposed strategy are affected by several factors. Such include the farming system, acceptability of both consumers and farmers, policies, and financial support [10].

As the majority of CH₄ is produced in the rumen, strategies for mitigating CH₄ emissions, necessarily affect the rumen microbiome and vice-versa. The rumen is harbored by a consortium of microorganisms including protozoa, bacteria, archaea, and fungi that conjoinly enable the fermentation of otherwise indigestible carbohydrates such as cellulose and hemicellulose into volatile fatty acids (VFA; e.g., acetate, propionate, and butyrate). Other products of carbohydrate fermentation include formate, ethanol, lactate, succinate, branched-chain volatile fatty acids, ammonia, CO₂, and H₂ [11]. Dissolved H₂ and CO₂ are utilized by methanogens, a group of microorganisms belonging to the Archaea domain, to form CH₄. Despite being the main pathway to avoid the accumulation of H₂ in the rumen, methanogenesis constitutes a loss of energy, reflected in animal productivity (e.g., methane emission has been correlated to residual feed intake [12]) and is strongly affected by diet (e.g., the level of concentrate has been correlated to CH₄ yield in grazing cows [13, 14]). However, methanogenesis is not the only H₂ sink mechanism in the rumen [15], being alternative pathways worth exploring. Therefore, the present work aims to summarize mitigation of CH₄ production, via alternative H₂ disposal pathways, namely, propionate production and reductive acetogenesis.

Microbial composition of the rumen

The rumen is a foregut ecosystem that hosts an enormous number of microbes living in symbiosis with the host. It is estimated to harbor a concentration of archaea of 10⁷ to 10⁸ cells/ml, bacteria of 10¹⁰ to 10¹¹ cells/ml, protozoa of 10⁴ to 10⁶ cells/ml, and fungi of 10³ to 10⁶ cells/ml [16]. Putting these into perspective, the number of ruminal microbial cells of an adult dairy cow (estimated volume of 50 to 200 L) [17] is 40 to 500 times the number of human
cells constituting an adult body (assuming the estimation of $3.72 \times 10^{13}$ total cells by Bianconi et al. [18]).

**Archaea**

Archaea in the rumen consist of methanogens belonging to the phylum Euryarchaeota [19]. It is represented by four orders Methanobacteriales, Methanococcales, Methanomicrobiales, and Methanosaurospirales [20]. Methanobacteriales include the genera Methanobacterium and the dominant genus Methanobrevibacter, which is divided into two clades, (1) clade Methanobrevibacter gottschalki that also includes the species Mbb. thaueri and Mbb. millerae; and (2) clade Methanobrevibacter ruminantium that also includes Mbb. olleyae [21]. Most methanogens use H$_2$ for the reduction of CO$_2$ into CH$_4$ [22], although formate might also be used instead of CO$_2$ by Methanobrevibacter, some strains of Methanobacterium spp., and Methanomicrobiales, such as genus Methanomicrobium [20]. In addition, methanogens of order Methanococcales and Methanosaurospirales can utilize methyl groups (e.g., genus Methanosphaera also uses methanol, whereas genera Methanosarcina and Methanomicromonas also use methylamines), and acetate (e.g., Methanosarcina) to produce CH$_4$ [20]. Methanol is originated from the demethylation of dietary pectins, whereas mono-, di-, and tri-methylamines are mainly end-products of plant phosphatidylcholine degradation [23]. The structure of the methanogenic community at the species or strain level has been correlated with feed efficiency [24]. Moreover, some authors argue that feed efficiency is related to CH$_4$ emissions [12, 25], although the link between both is complex and influenced by multiple parameters related to the rumen microbiome [26] and host factors (e.g., passage rate and nutrient absorption) [27]. Importantly, dominant archaea groups were found similar in samples collected from ruminants across the globe, which is likely an advantage to develop and implement worldwide strategies to mitigate CH$_4$ emissions targeting methanogens [19].

**Bacteria**

The rumen harbors cellulolytic and non-cellulolytic bacteria, being the first able to degrade cellulose and hemicelluloses. Primary cellulose fermenters are Fibrobacter succinogenes, Ruminococcus flavefaciens, and Ruminococcus albus. These bacteria are non-motile, adhering exclusively to the fibers through the glycocalyx, and having cellulosomes located on the cell surface [28]. They hydrolyze cellulose and other polysaccharides (e.g., hemicelluloses and pectin), producing celldextrins to utilize as a source of energy and make available for cross-feeding [28]. This is important to provide nutrients for the growth of other bacteria and/or non-adherent cells of the same species that are poised to adhesion to new feed particles [29]. Secondary cellulose fermenters, including Butyrivibrio fibrisolvens, Clostridium longisporum, and Clostridium l. leodaeidii, might be motile or non-motile, adhering minimally to fibers, and having extracellular cellulases [28]. Non-cellulolytic bacteria, able to degrade starch, hemicelluloses, or pectin, might include Prevotella ruminantium, Eubacterium xylanophilum, Ruminobacter amylophilus, Succinimonas amylolytica, Succinivibrio dextrinosolvens, Selenomonas ruminantium, Selenomonas lactilytica, Lachnospira multiparas, Streptococcus bovis, and Megasphaera elsdenii [30].

Despite the great diversity of bacterial species in the rumen, an extensive study reported the existence of 30 most abundant bacterial groups comprising $\approx 89\%$ of total sequences found on livestock species [19]. Among them, Prevotella, Butyrivibrio, and Ruminococcus, unclassified Lachnospiraceae, Ruminococcaceae, Bacteroidales, and Clostridiales were predominant, having an abundance of $\approx 67\%$. Later works analyzing metagenome-assembled genomes (MAGs) revealed the existence of new genomes from the Actinobacteria, Fibrobacteres, and Proteobacteria phyla, also highlighting the abundance of genus Succinivibrio [31].

The existence of a core ruminal bacterial microbiome has been suggested, despite a clear variation associated with host and diet [19]. Indeed, a study reported that the breed is more determinant for metabolites and bacterial communities of bovines than diet and life-stage [32]. Even though the age and time of weaning significantly affect the diversity and abundance of rumen microbial communities [33], it has been suggested that colonization of rumen starts in utero, as samples collected in goat fetuses allowed the identification of sequences mainly belonging to the phylum Proteobacteria [34]. Also, three moments were shown to produce important shifts in rumen bacteria: delivery, milk intake, and weaning [34]. Moreover, it was reported that the rumen microbial colonization is affected differently by natural or artificial milk feeding systems [35], and the inoculation of young ruminants with fresh rumen fluid from adult animals enhanced microbial colonization, likely improving rumen development [36]. Furthermore, early life modulation of the rumen microbiome has been a matter of study regarding its effectiveness when compared to later interventions for the improvement of rumen fermentation and reduction of CH$_4$ emission [37].

**Protozoa**

Contrary to archaea and bacteria, ciliate protozoa vary across ruminants of different species as well as individuals of the same species [19]. A study in cattle indicated that Entodinium, Diplodinium, Eremoplastron,
Ostracodinium, Eodinium, Epidinium, Isotricha, and Dasytricha were among the most abundant genera (>1%) of a total of 13 identified in the ruminal fluid by microscopic identification and counting [38]. Conversely, in sheep and goats, 12 and 8 genera were identified, being Dasytricha, Entodinium, Eudiplodinium, Diplodinium, Isotricha, and Metadinium the more abundant (>1%) in both species, and Enoploplastron, Ophryoscolex, and Polyplastron only in sheep [39].

Protozoa attach to the surface of partially digested feed particles, in which other microorganisms exert high fibrolytic activity, allowing them to take advantage of monosaccharides (e.g., glucose, cellobiose, and celodextrins) that they use as their source of energy for growth and metabolism [40]. Also, protozoa predate bacteria from which they obtain acids for growth and maintenance [41]. Non-surprisingly, ciliate protozoa affect the diversity of ruminal bacteria [42] and end-fermentation products. However, protozoa are not essential for ruminal fermentation and removal of protozoa from the rumen, also called defaunation, does not seem to significantly impact animal health, although feed digestibility might be affected [41]. The potential of defaunation to mitigate CH4 emission has been studied [43, 44]. The external surface of protozoa is a site for methanogen attachment (ectosymbionts) or intracellular colonization (endosymbionts), which are attracted by the H2 produced in protozoan hydrogenosomes [41]. This symbiosis enhances methanogenesis and favors protozoa as it reduces the levels of H2, enabling them to continue the fermentation of monosaccharides left by bacteria. Indeed, the protozoan-associated methanogens are estimated to be responsible for 37% of CH4 emission [45], and a meta-analysis including several in vivo studies concluded that a reduction of protozoa concentration was, in most cases, associated with a reduction of CH4 emission [46]. Nevertheless, as noted by some authors, defaunation has implications on other metabolic pathways (e.g., fatty acids), which likely justifies a holistic view of the use of this strategy for CH4 mitigation [47]. Ionophore additives were suggested for reduction of CH4 emission, due to their positive [48] and transient effects [49] on reduction of protozoans and methanogens. However, environmental contamination with still active antibiotics is detrimental for the environment and needs to be considered [50]. Other strategies targeting rumen protozoans include the supplementation of plant metabolites, such as essential oils, which disrupt protozoal membrane, indirectly reducing methanogens [51–53].

Fungi
More than 90% of fungi sequences isolated in the rumen remain unclassified [54], however, the presence of anaerobic fungi belonging to phylum Neocallimastigomyces (genera Neocallimastix, Caecomyces, Pirromyces, Anaeromyces, Orpinomyces, and Cyllamyces) is acknowledged [55]. Fungi are infrequently found in strained rumen fluid because zoospores attach and colonize (encyst and germinate to produce the fungal thallus) the plant fragments suspended in the rumen natural digesta, being only then released by the sporangia [56]. This may have contributed to a greater unawareness of the importance of their fibrolytic activity. Nonetheless, fungi have enzymes to degrade plant cell wall carbohydrates [55] and are indeed more efficient in degrading lignin than bacteria [57]. In co-culture with methanogens, Neocallimastix exhibited high lignocellulose-degrading activity with the production of CH4 and acetate [58]. Other studies support the importance of fungi as substrate and electron donors for methanogenesis [59], which makes fungal metabolic pathways an attractive subject for study in the context of the mitigation of CH4 emission.

Ruminal carbohydrate digestion
Plant cell wall polysaccharides are arrangements of glycosidic linkages (e.g., mono-, di-, and oligosaccharides) and noncarbohydrate moieties [60], and might be analytically grouped into cellulose, hemicellulose, and pectin [30]. Cellulose, the most abundant component of cell wall plants, is formed by β-glucose and other hexoses. Whereas, hemicellulose, mainly composed of pentoses with linear xylose chains and variable linkages of arabinose, uronic acids, and galactose, is the second most abundant [61]. Pectin is present in the primary cell wall and has D-galacturonate in its structure [30]. Moreover, starch, a non-structural carbohydrate, is composed of α-glucose, constituting the major carbohydrate storage in plants and an important source of energy for ruminants [62]. Ruminal microorganisms cleave complex glycosidic bonds mainly through glycoside hydrolases. Indeed, the enrichment in those enzymes observed in Bacteroidales, including Prevotellaceae, Fibrobacteres, and some Clostridiales [63], affords them a competitive advantage justifying its higher abundance in the rumen.

Chewing and rumination are important for carbohydrate digestion carried out by ruminal microorganisms, as it facilitates the adhesion of bacteria to plants, hydrating and disrupting the protective cuticular layer of plants [29]. After specific adhesion, in which bacterial-substrate linkages and adhesins are developed, the proliferation and colonization of plant tissues are initiated [29]. This is possible because the fermentation of sugars leads to the formation of ATP, the main source of energy for microorganisms [61]. Microbial enzymes degrade the hexoses primarily through the Embden-Meyerhof pathway, originating NADH and pyruvate. Hemicellulose
has two pathways for degradation, the transketolase and transaldolase reactions of the pentose cycle or phosphoketolase, with the products originated entering the Embden-Meyerhof pathway [61]. Continuation of the metabolism of the intermediate compound pyruvate is dependent on the oxidation of cofactors (e.g., NADH) through pathways that lead to the formation of lactate, succinate, acetate, propionate, butyrate, ethanol, and valerate [64]. The several reactions that occur in rumen fermentation entail the formation and incorporation of metabolic hydrogen [H]. Formation corresponds to the transfer of electron donors of metabolic intermediates to oxidized intracellular cofactors (highlighted in pink in Fig. 2). Conversely, incorporation corresponds to the transfer from reduced intracellular cofactors to metabolic intermediate electron acceptors (highlighted in green in Fig. 2) [65]. This electron transfer is carried out by hydrogenases and originates H₂. Furthermore, H₂ and CO₂ might also be originated from the conversion of formate in the pyruvate-ferredoxin oxidoreductase [64]. Hydrogen is then transferred between producing species (bacteria, protozoa, and fungi) and hydrogenotrophic microorganisms, mainly methanogens.

**Methanogenesis and CH₄ emission**

Rumen methanogenesis is known to occur by three different pathways: hydrogenotrophic (A), acetoclastic (B), and methylotrophic (C) [20]:

(A) \(4H_2 + CO_2 \rightarrow CH_4 + 2H_2O\)
\(4HCOOH \rightarrow CH_4 + 3CO_2 + 2H_2O\)

(B) \(CH_3COOH \rightarrow CH_4 + CO_2\)

(C) \(4CH_3OH \rightarrow 3CH_4 + CO_2 + 2H_2O\)
\(4CH_3 = NH_2 + 2H_2O \rightarrow 3CH_4 + CO_2 + 4NH_3\)
\(2(CH_3)_2 = NH + 2H_2O \rightarrow 3CH_4 + CO_2 + 2NH_3\)
\(4(CH_3)3 = N + 6H_2O \rightarrow 9CH_4 + 3CO_2 + 4NH_3\)

Hydrogenotrophic methanogenesis is largely the most frequent, which likely occurs because the energetics involved is more favorable, resulting in slower growth rates and lower cell yields for microorganisms involved in the other methanogenesis pathways [22]. The genus *Methanobrevibacter*, a highly abundant hydrogenotrophic, has been correlated with high CH₄ emissions in steers [66], heifers [67], and dairy cows (more particularly *Mbb. gottschalkii* and *Mbb. ruminantium*) [68]. In turn, *Methanosphaera* (methylotroph) was negatively correlated with CH₄ emission in heifers [67] and dairy cows [69, 70]. This might be explained by the stoichiometric of the reaction, as one mole of CO₂ is required to produce one mole of CH₄ (A), while four moles of methanol are required to produce three moles of CH₄ (B) in the methylotrophic pathway [69]. Interestingly, the methylotrophic pathway appears to be more significant in young calves compared to mature cows [71]. Even though methylotrophs might have a lower CH₄ yield, they release NH₃ to add to the amount already produced by proteolytic bacteria [72]. Importantly, methyl-coenzyme M reductase, which catalyzes the final step of methanogenesis is common across the different methanogenesis pathways [73], which is likely an advantage for strategies intending to target directly methanogens. An example is 3-nitroxypropional, a molecule that was shown to inhibit methanogenesis by oxidizing the active site Ni(I) of methyl-coenzyme M reductase [74].

Despite the differences reported in microbial abundances, metagenomic and metatranscriptomic sequencing studies showed that the increased expression of methanogenesis pathway genes explains the increase in CH₄ emissions, which might itself be regulated by a substrate effect [75]. The diversity of methanogens in the rumen has been correlated to CH₄ emission. A study using co-abundance analysis of rumen microorganisms of cows reported that low-CH₄ emitting animals had a more diverse community of methanogens involved in the three methanogenic pathways, compared with high-CH₄ emitting ones, which had low numbers of hydrogenotrophic methanogenic genera [76]. This shows that methanogen diversity is correlated to CH₄ emission and highlights the interaction between communities and competition among methanogens for H₂. Also, low-CH₄ cow emitters exhibited a more complex microbial network composed of a diverse microbiome, more specifically, bacterial and fungal genera and their genes [76]. Indeed, differences in the microbiome of low- and high-CH₄ emitters are not limited to methanogens, which is not surprising as the availability of precursors for methanogenesis is dictated by the fermentative microbial consortium. Experimentally, *S. dextrinosolvens* in co-culture enhanced a member of Methanomassiliicoccales and inhibited the activity of a member of the *Mbb. gottschalkii* clade [77]. An in vivo study showed that low-CH₄ emitting sheep were associated with an elevated abundance of l-lactate dehydrogenase genes, an enrichment of genus *Sharpea* (family *Erysipelotrichiaceae*), and decreased abundance of families *Lachnospiraceae* and *Ruminococcaceae* [78]. These results were confirmed by another study using MAG on microbial sequences collected in high and low CH₄ emitting sheep, allowing the identification of differential abundance of other genera and microorganisms at the species level [31]. Low-CH₄ emitting sheep had, in addition to previously mentioned, a high abundance of *Kandleria*, *Fibrobacter*, and *Selenomonas* at a genus level, whereas at a species level, *Fibrobacter succinogenes* and several species of *Bifidobacterium*, *Olsenella*, *Desulfovirio* were more abundant [31], compared to high emitting sheep. In dairy cows, the abundance of genera...
Fig. 2 Scheme of rumen carbohydrate fermentation pathways into volatile fatty acids (bold lined boxes) and other intermediate metabolites. The rumen H₂ sink pathways are displayed: reductive acetogenesis (1), methanogenesis (2), sulfate reducers (3), and nitrate reducers (4).
Christensenellaceae, Mogibacteriaceae, S24-7, Butyricibrio, Shwartzia, and Treponema was associated with low CH₄ emission in one study [69] and with genus Eubacterium in another, in which rumen samples of heifers were analyzed [67]. It is important to consider that effects of the host (e.g., ruminant species), diet as well as methodologies (e.g., sequencing technique and/or 16S rRNA hypervariable region selected) might preclude a direct comparison of microbial composition across studies.

Age and physiological states affect the rumen microbiome composition and thus CH₄ yields, as shown in a study in heifers, in which the interaction of genera Prevotella and Methanobrevibacter was associated with the CH₄ yield, while in older cows, the significant interaction was between Methanobrevibacter and Succinivibrio [79]. Interestingly though, a study reported that early and late lactation stages were correlated with different levels of CH₄ emissions not accompanied by changes in the rumen microbiome [80]. In a study with Colombian buffalos, the genus Prevotella was associated with low CH₄ emissions [81]. Despite the differences reported in archaea composition between buffalo and cattle [82], this bacterial group seems relevant for CH₄ emission of both species. As some authors pointed, the host genotype affects the phenotype of CH₄ emission, in addition to its microbial community [81]. Indeed, one study indicated that H₂-producing bacteria explained up to 24% of CH₄ phenotypic variance, and host genome, 14% [70]. In another study, the CH₄ emission had a cumulative effect of archaea and bacteria of 13% and 21% of host genetics [83]. Despite the percentage difference, both studies suggest that targeting the rumen microbiome of low-CH₄ emitting animals (through breeding programs) is possible and likely more effective than not considering the host genome and heritability of the trait.

**Alternative pathways to H₂ sink**

As previously stressed, the concentration of H₂ determines the CH₄ produced, while the production of H₂ is determined by the prevailing pathways of glucose fermentation. The fermentation into butyrate (D) and acetate (E) entails a potential production of one mole of H₂ (per mole of glucose), whereas propionate (F) entails a net incorporation of one mole of H₂ (per mole of glucose) [65]. This balance considers the reducing equivalents [2H⁺] produced and incorporated through several reactions (Fig. 2). Microbial cells able to change their fermentation patterns and better adapt to certain conditions are likely more active degrading the available substrates, thus managing to expand [84]. The concentration of H₂ conditions the fermentation pathway, affecting the free energy change between reactants and products in which the microbial biomass thrives. Summarily, high concentrations of H₂ favor the productions of propionate, whereas low concentrations of H₂ favors the production of acetate [11]:

(D) C₆H₁₂O₆ → 2CH₃CH₂CH₂COO⁻ + CO₂ + 2H₂ + H⁺
(E) C₆H₁₂O₆ + 2H₂O → 2CH₃COO⁻ + 2CO₂ + 2H⁺ + 4H₂
(F) C₆H₁₂O₆ → CH₃CH₂COO⁻ + 2H₂O + 2H⁺

A recent study analyzing MAGs of the microbiota of gastrointestinal ruminants reported that 48% encoded enzymes for fermentative H₂ production, 1.5% for H₂-uptake hydrogenases and the methyl-coenzyme M reductase (mcrA genes) related to hydrogenotrophic methanogenesis, while 11% encoded both hydrogenases and the required terminal reductases of alternative methanogenesis pathways. Acetogenesis constituted 3% of MAGs, furamurate 1.9%, and sulfate reduction 0.8% [26]. It has been proposed that the microbial hydrogenases and fermentation pathways are differentially regulated through direct H₂ sensing by putative sensory [FeFe]-hydrogenases [15]. Indeed, sensory hydrogenases, as well as fermentative and bifurcating hydrogenases, are highly expressed in Clostridiales, Bacteroidales, and Selenomonadiales, whereas methanogenic hydrogenases are present, by order of expression, in Methanobacteriales > Methanomassiliicoccales > Methanosarcinales [15]. Moreover, low-CH₄ emitting sheep were reported to have higher hydrogenase and terminal reductase transcripts from alternative H₂ uptake pathways, which might even serve as a larger H₂ sink than methanogenesis, compared to high-CH₄ emitters [15]. This highlights the preponderant role of the bacterial consortium in determining H₂ metabolism and the strategies of methanogens to compete and affect H₂ utilization, and thus CH₄ emission.

Dietary manipulation of H₂ production has been attempted to reduce CH₄ emissions, and it is highly associated with changes in ruminal microbiota. Higher concentrations of H₂ are associated with high starch content diets, likely because H₂ release outgrows the capacity of H₂-consuming microorganisms, leading to an accumulation [85]. Hydrogen accumulation was also associated with diets containing tannin-rich peanut skin provided to beef cattle due to the reduction of H₂-using microorganisms, including populations of Bacteroidetes phylum, total methanogens, Methanobrevibacter, and protozoa, concomitantly reducing methanogenesis [84]. Supplementation of non-fermentative sources, such as Mg, was shown to increase ruminal dissolved H₂, which affected the microbiota, decreasing the copy number of fungi in goats [86].

In addition to diet effects, host factors and H₂-producing microorganisms interact, conditioning H₂ production. Indeed, a study in sheep reported that
animals with a smaller rumen volume emitted proportionally less CH₄, presumably because their higher feed rate passage selected microorganisms able to grow on soluble sugars that can be quickly degraded [78], thus increasing H₂.

Directly targeting methanogens and potentiating other H₂-consuming microorganisms (e.g., propionate pathway, reductive acetogenesis, nitrate, and sulfate reduction) have been proposed as strategies to mitigate CH₄ emissions. Sulfate-reducing bacteria (e.g., nitrate-reducing propionibacteria, Wolinella succinogenes, and Veillonella parvula) can reduce nitrate and nitrocompounds into N₂O and NH₄ [87], and although studies [88] have proven it effective in decreasing methanogenesis, concerns over toxicity have hindered its use and motivated further research to develop safer additives (e.g., encapsulated nitrite [89]). Sulfate-reducing bacteria (e.g., Desulfovibrio) reduce sulfate into H₂S, competing with methanogens for H₂ and reducing methanogenesis if ruminal levels of sulfate increase [90]. Despite thermodynamics and matrix affinity favors sulfate reduction over methanogenesis, as seen in marine sediments [90, 91], the product is highly toxic to the animal.

Propionate pathway

Ruminal propionate originates mainly from succinate and acrylate pathways. The fermentation of carbohydrate-rich diets with high levels of soluble sugars promotes the proliferation of amylolytic microorganisms such as S. bovis [92], Lactobacillus, and Bifidobacterium [93] that reduce pyruvate into lactate. In physiological conditions (no acidosis), microorganisms such as M. elsdenii and Coprococcus catus produce propionate from lactate via the acryl-CoA, using the acrylate pathway [94]. Indeed, both M. elsdenii and C. catus were correlated to higher feed efficiency and lower CH₄ emissions in dairy cows [95].

The production of propionate from lactate can also occur with a first oxidation to pyruvate, followed by carboxylation to oxaloacetate, reduction to malate, dehydration to fumarate, reduction to succinate, and a final decarboxylation to propionate (Fig. 2) [96]. This, also called the ‘randomizing’ pathway, might be carried out by microorganisms such as S. ruminantium [97] and Succinilbacterium ruminis (unable to ferment substrates other than succinate) [98]. According to an in vitro study, the addition of fumarate-reducing bacteria, Mitsuokella jalaludinii, lowered methanogen DNA copies and occurrence through competition for H₂ [99]. Similarly, supplementation with enterococci (E. faecalis and E. faecium) exhibited fumarate reductase activity, with an increase of propionate and a decrease of CH₄ [100]. Another in vitro study reported the potential effects of propionic bacteria on ruminal feed degradation by showing a reduction of CH₄ from 8 to 20% associated with one strain of Propionibacterium jensenii and two of Propionibacterium thoenii [101]. Lactiplantibacillus plantarum was also shown to decrease in vitro CH₄ production and increased propionate [102]. In turn, an in vivo study showed that, when provided for 4 weeks to lactating primiparous cows fed contrasting high-starch or high-fiber diets, Propionibacterium freudenreichii, Lactiplantibacillus pentosus, and Lactobacillus delbrueckii subsp. bulgaricus did not affect CH₄ emissions [103]. This highlights the need to further study the in vivo effects of lactic acid bacteria to elucidate any potential correlation with the in vitro positive results. The use of lactic acid bacteria requires finding delivery options that have already been implemented in the global farming system (e.g., silage inoculants and direct-fed microbes), a challenge that must be addressed [104].

Manipulation of fiber content, namely by the replacement of forage fiber by non-forage fiber sources, promoted an expansion of Firmicutes over Bacteroidetes and of Methanobrevibacter over Methanomassiliicoccus, and a successful shift of H₂ flow towards the propionate pathway [105]. Furthermore, feed supplementation with saponin (which causes cell rupture and lysis of protozoan) was shown to shift fermentation products, lowering butyrate and increasing propionate [106].

In vivo studies associated other bacteria such as genera Succinivibrio (family Succinivibrionaceae), Roseburia, and Blautia (family Lachnospiraceae) with the increase of propionate, when testing diets that differed in the corn processing techniques [107]. Furthermore, family Succinivibrionaceae were effectively associated with high propionate and low CH₄ yield [66, 108].

Adding to its potential as a methanogenesis competitor, the increase of propionate production is advantageous for cows’ health and efficiency. The major share (50 – 75%) of propionate produced is absorbed into the portal vein [109]. Propionate is the main precursor of hepatic gluconeogenesis and essential to supply glucose to the mammary gland, thus contributing to support milk production [110]. Furthermore, infusions of propionate led to greater plasma progesterone concentrations post-ovulation, which can affect follicular development and pregnancy rates, and thus improve reproductive efficiency [111].

Reductive acetogenesis

Acetogenesis has been documented as an alternative to methanogenesis, in which via the acetyl-CoA pathway, two moles of CO₂ and four moles of H₂ are incorporated per one mole of acetate produced (G), in a thermodynamically feasible set of reactions.
Acetogens degrade multiple substrates (e.g., pentoses, hexoses, alcohols, formate, and methyl groups) in addition to $\text{H}_2$, and in the rumen of cows and sheep, the dominant microorganisms are members of Lachnospiraceae, Clostridiaceae, and Ruminococcaceae families [112, 113]. The threshold value for $\text{H}_2$ utilization is lower for methanogens than it is for acetogens, which renders reductive acetogenesis a disadvantage. Therefore, increasing $\text{H}_2$ pressure would remove methanogenesis thermodynamic advantage [114]. An in vitro study reported that supplementation of acetogen Eubacterium limosum was able to produce acetate when methanogens were suppressed concomitantly, but a minimal change in $\text{CH}_4$ production was observed when methanogenesis was not inhibited [115]. Similarly, in vitro supplementation with E. limosum and Proteiniphilum acetatigenes decreased $\text{CH}_4$ concentrations, increased acetate, whereas in vivo, P. acetatigenes was associated with high milk protein, lower somatic cell counts, and lower decline of milk production over 60 days [116].

Conclusions
Several in vivo studies employing sequencing techniques have revealed distinct ruminal microbial communities between high- and low-$\text{CH}_4$ emitting animals. That knowledge has recently been prompted by metagenomics and transcriptomics that allow not only studying the rumen microbial composition but also its function, unraveling genes and pathways involved in the metabolism of $\text{H}_2$. The goal of successful interventions is likely shifting $\text{H}_2$ from methanogenesis, exploiting natural processes, without compromising rumen physiology. That depends upon the interaction between microbial communities, including several orders of bacteria, archaea, and eukaryotes that encode and express enzymes [15], thus mediating ruminal fermentation pathways. The propionate pathway competes with methanogenesis, improving efficiency. Using additives including probiotics to promote propionate production showed good results in vitro, yet in vivo studies are still required to confirm its efficacy as well as its suitability for widespread use. Dietary management, namely the use of concentrates and high-rich-carbohydrate diets that naturally select microorganisms involved in the propionate pathway make this a more suitable strategy for intensive production systems. However, the carbon footprint of diet production and transportation must be placed in the equation. Reductive acetogenesis is suitable for grazing animals, in which degradation of fibrous diets originates $\text{H}_2$ that can be utilized by acetogens, though a combined strategy to reduce methanogens is required.

Even though knowledge on the ruminal microbiome has expanded, there are still several genes and microorganisms whose characterization and function remains unknown, particularly protozoan and fungi. Recent works have recovered archael and bacterial MAGs from metagenomic data, from which new, previously unnoticed, pathways and networks have been discovered. This indicates that the ruminal microbiome study is a work in progress and may provide us new prospects for finding solutions for lowering livestock enteric $\text{CH}_4$ emissions, thus addressing the role of cattle in the current climate emergency.
et al. Animal Microbiome

5. Gougoulias C, Clark JM, Shaw LJ. The role of soil microbes in the global carbon cycle: tracking the below-ground microbial processing of plant-derived carbon for manipulating carbon dynamics in agricultural systems. J Sci Food Agric. 2014;94:2362–71.

6. Raiten DJ, Allen LH, Slavin JL, Mitloehner FM, Thoma GJ, Haggerty PA, et al. Understanding the intersection of climate/environmental change, health, agriculture, and improved nutrition: a case study on micronutrient nutrition and animal source foods. Curr Dev Nutr. 2020;4:1–8.

7. González‑Recio O, López‑Paredes J, Ouatahar L, Charfeddine N, Ugarte CR, Serrano‑Gaona VH, Calzada‑Chavez MP, Camacho‑García M, Crespo‑Ramírez C, Corella‑Rodríguez MP, et al. Mitigation of greenhouse gases in dairy cattle via genetic selection. 2. Incorporating methane emissions into the breeding goal. J Dairy Sci. 2020;103:7210–21. https://doi.org/10.3168/jds.2019‑19758.

8. Baca‑González V, Asensio‑Calavia P, González‑Acosta S, de la Lastra JMP, de la Nuez AM. Are vaccines the solution for methane emissions from ruminants? A systematic review. Vaccines. 2020;8:1–21.

9. Haque MN. Dietary manipulation: a sustainable way to mitigate methane emissions from ruminants. J Anim Sci Technol. 2018;60.15. https://doi.org/10.1186/s40781‑018‑0175‑7.

10. Key N, Tallard G. Mitigating methane emissions from livestock: a global analysis of sectoral policies. Clim Change. 2012;112:387–414. https://doi.org/10.1007/s10584‑011‑0206‑6.

11. Janssen PH. Influence of hydrogen on rumen methane formation and fermentation balances through microbial growth kinetics and fermentation thermodynamics. Anim Feed Sci Technol. 2010;160:1–22.

12. Fitzsimons C, Kenny DA, Deighton MH, Fahey AG, Mc Gee M. Methane emissions, body composition, and rumen fermentation traits of beef heifers differing in residual feed intake. J Anim Sci. 2013;91:5789–800. https://doi.org/10.2527/jas.2013‑6956.

13. Jiao HP, Dale AJ, Carson AF, Murray S, Gordon AW, Ferris CP. Effect of concentrate feed level on methane emissions from grazing dairy cows. J Dairy Sci. 2014;97:7043–53. https://doi.org/10.3168/jds.2014‑7979.

14. van Wyngaard JDV, Meeske R, Erasmus LJ. Effect of concentrate level on enteric methane emissions, production performance, and rumen fermentation of Jersey cows grazing kikuyu‑dominant pasture during summer. J Dairy Sci. 2018;101:9954–66. https://doi.org/10.3168/jds.2017‑14327.

15. Greening C, Geier R, Wang C, Woods LC, Morales SE, McDonald MJ, et al. Diverse hydrogen production and consumption pathways influence methane production in ruminants. ISME J. 2019;13:2617–32.

16. Wright ADG, Klieve AV. Does the complexity of the rumen micro‑biological ecology preclude methane mitigation? Anim Feed Sci Technol. 2011;166–167:248–53.

17. Membrive CMB. Anatomy and physiology of the Rumen. In: Millen DD, De Beni AM, Lauritano Pacheco RD, editors. Rumenology. 1st ed. Cham: De Beni AM; 2012. p. 1–34. https://doi.org/10.1007/978‑3‑ 319‑ 30533‑2_1.

18. Bani coni E, Piovesan A, Facchin F, Beraudi A, Casadei R, Frabetti F, et al. An estimation of the number of cells in the human body. Ann Hum Biol. 2013;40:463‑71. https://doi.org/10.3109/03014460.2013.807787.

19. Rada L, Weyler DR, Alberts CW, McEvoy CJ, Bäuerle MA, et al. Rumen microbial community composition varies with diet and host, but a core microbiome is found across a wide geographical range. Science. 2018;361:658–62. https://doi.org/10.1126/science.aar3710.

20. Liu Y, Whitman WB. Metabolic, phylogenetic, and ecological diversity of the methanogenic archaea. Ann N Y Acad Sci. 2008;1125:171–89.

21. Janssen PH, Kirs M. Structure of the archaean community of the rumen. Appl Environ Microbiol. 2008;74:3619–25.

22. Fenchel T, King GM, Blackburn TH. Bacterial Metabolism. In: Fenchel T, King GM, Blackburn T‑HT, editors. Bacterial biogeochemistry. 3rd ed. Boston: Academic Press; 2012. p. 1–134. https://doi.org/10.1007/978‑0‑12‑415836‑8.00001‑3.

23. Kelly WJ, Leahy SC, Kamke J, Soni P, Koike S, Mackie R, et al. Occurrence and expression of genes encoding methyl‑methyl compound production in rumen bacteria. Anim Microbiome. 2019;1:1–13.

24. Zhou M, Hernandez‑Sanabria E, Guan LL. Characterization of variation in rumen methanogenic communities under different dietary and host feed efficiency conditions, as determined by PCR‑Denaturing Gradient Gel Electrophoresis analysis. Appl Environ Microbiol. 2010;76:3776‑86. https://doi.org/10.1128/AEM.00010‑10.

25. Sakamoto LS, Souza LL, Gianvecchio SB, de Oliveira MHV, Silva JV, Cane sin RC, et al. Phenotypic association among performance, feed efficiency and methane emission traits in Nellore cattle. PLoS ONE. 2021;16:e0257964. https://doi.org/10.1371/journal.pone.0257964.

26. Ozutsumi Y, Tajima K, Takenaka A, Ibarashy H. The effect of protozoa on the composition of rumen bacteria in cattle using 16S rRNA gene clone libraries. Biosci Biotechnol Biochem. 2003;67:499‑506.

27. Palma‑Hidalgo JM, Jiménez E, Popova M, Morgavi DP, Martín‑García AM, de Beni AM, Lauritano Pacheco RD, editors. Rumenology. 1st ed. Cham: De Beni AM; 2012. p. 1–34. https://doi.org/10.1007/978‑3‑ 319‑ 30533‑2_1.

28. Zhou M, Hernandez‑Sanabria E, Guan LL. Characterization of variation in rumen methanogenic communities under different dietary and host feed efficiency conditions, as determined by PCR‑Denaturing Gradient Gel Electrophoresis analysis. Appl Environ Microbiol. 2010;76:3776‑86. https://doi.org/10.1128/AEM.00010‑10.

29. Sakamoto LS, Souza LL, Gianvecchio SB, de Oliveira MHV, Silva JV, Cane sin RC, et al. Phenotypic association among performance, feed efficiency and methane emission traits in Nellore cattle. PLoS ONE. 2021;16:e0257964. https://doi.org/10.1371/journal.pone.0257964.

30. Wang Y, McAllister TA. Rumen microbes, enzymes and feed digestion: A Review. Asian‑Australas J Anim Sci. 2002;15:1659‑74. https://doi.org/10.1017/10584‑011‑0206‑6.
Belanche A, De La Fuente G, Newbold CJ. Effect of progressive inocula-

Guyader J, Eugène M, Nozière P, Morgavi DP, Doreau M, Martin C. Influence of rumen protozoa on methane emissions in ruminants: a meta-analysis approach. Animal. 2014;18:1621–56.

Francisco AE, Santos-Silva JM, Portugal APV, Alves SP, Bessa RJB. Relationship between rumen ciliate protozoa and biohydrogenation fatty acid profile in rumen and meat of lambs. PLoS ONE. 2019;14:1–21.

Guan H, Wittenberg KM, Ominski KH, Krause DO. Efficacy of iono-

Crossland WL, Tedeschi LO, Callaway TR, Miller MD, Smith WB, Cravey M. In vitro effects of rotating antibiotic and ionophore feed additives on volatile fatty acid production, potential for methane production, and microbial populations of steers consuming a moderate-forage diet. J Anim Sci. 2017;95:4545–67. https://doi.org/10.2527/jas.2017.1665.

Takahashi J, Iwasa M. Entomological approach to the impact of ionophore-feed additives on greenhouse gas emissions from pasture land in cattle. J Anim Sci Technol. 2021;63:16–24. https://doi.org/10.5187/jast.2021.e11.

Ku-Verá JC, Jiménez-Ocampo R, Valencia-Salazar SS, Montoya-Flores MD, Molina-Botero IC, Arango J, et al. Role of secondary plant metabo-

Janusz G, Pawlik A, Sulej J, Świderska-Burek U, Jarosz-Wilkołazka A, et al. The rumen microbial metagenome associated with high methane production in cattle. FEMS Microbiol Ecol. 2015. https://doi.org/10.1093/femsec/fiu026.

Russell JB, Wallace RJ. Energy-yielding and energy-consuming reactions. In: Hobson PN, Stewart CS (editors). The rumen microbial ecosys-

Ungerfeld EM. Metabolic hydrogen flows in rumen fermentation: prin-
ciples and possibilities of interventions. Front Microbiol. 2020;11:589.

Wallace RJ, Rooke JA, McKain N, Dutthie CA, Hyslop JJ, Ross DW, et al. The rumen microbial metagenome associated with high methane production in cattle. BMC Genomics. 2015;16:1–14.

Cunha CS, Marcondes MI, Veloso CM, Mantovani HC, Luiz B, Pereira GR, et al. Compositional and structural dynamics of the ruminal microbiota in dairy heifers and its relationship to methane production. Sci Food Agric. 2018;99:210–8. https://doi.org/10.1002/jsfa.9162.

Rothwell JP. The rumen microbial metagenome following the introduction of complex rumen microbiome interactors. Front Microbiol. 2020;11:583.

Friedman N, Jami E, Mizrahi I. Compositional and functional dynam-
ics of the bovine rumen microecosystem across different development stages. Environ Microbiol. 2017;19:3365–73.

Poulsen M, Schwab C, Borg Jensen B, Engberg RM, Søergaard N, et al. Methanogenic and bacterial community structure. Front Microbiol. 2017;8:226.

Cunha CS, Veloso CM, Marcondes MI, Mantovani HC, Tomich TR, Pereira LGR, et al. Assessing the impact of rumen microbial communities on methane emissions and production traits in Holstein cows in a tropical climate. Syst Appl Microbiol. 2017;40:492–9.

Ramayo-Caldas Y, Zingaretti L, Popova M, Estellé J, Bernard A, Pons N, et al. Identification of rumen microbial biomarkers linked to methane emission in Holstein dairy cows. J Anim Breed Genet. 2020;137:49–59.

Friedman N, Jami E, Mizrahi I. Compositional and functional dynam-
ics of the bovine rumen microecosystem across different development stages. Environ Microbiol. 2017;19:3365–73.

Poulsen M, Schwab C, Borg Jensen B, Engberg RM, Søergaard N, et al. Methanogenic and bacterial community structure. Front Microbiol. 2017;8:226.

Cunha CS, Veloso CM, Marcondes MI, Mantovani HC, Tomich TR, Pereira LGR, et al. Assessing the impact of rumen microbial communities on methane emissions and production traits in Holstein cows in a tropical climate. Syst Appl Microbiol. 2017;40:492–9.

Ramayo-Caldas Y, Zingaretti L, Popova M, Estellé J, Bernard A, Pons N, et al. Identification of rumen microbial biomarkers linked to methane emission in Holstein dairy cows. J Anim Breed Genet. 2020;137:49–59.

Friedman N, Jami E, Mizrahi I. Compositional and functional dynam-
ics of the bovine rumen microecosystem across different development stages. Environ Microbiol. 2017;19:3365–73.

Poulsen M, Schwab C, Borg Jensen B, Engberg RM, Søergaard N, et al. Methanogenic and bacterial community structure. Front Microbiol. 2017;8:226.

Cunha CS, Veloso CM, Marcondes MI, Mantovani HC, Tomich TR, Pereira LGR, et al. Assessing the impact of rumen microbial communities on methane emissions and production traits in Holstein cows in a tropical climate. Syst Appl Microbiol. 2017;40:492–9.

Ramayo-Caldas Y, Zingaretti L, Popova M, Estellé J, Bernard A, Pons N, et al. Identification of rumen microbial biomarkers linked to methane emission in Holstein dairy cows. J Anim Breed Genet. 2020;137:49–59.

Friedman N, Jami E, Mizrahi I. Compositional and functional dynam-
ics of the bovine rumen microecosystem across different development stages. Environ Microbiol. 2017;19:3365–73.

Poulsen M, Schwab C, Borg Jensen B, Engberg RM, Søergaard N, et al. Methanogenic and bacterial community structure. Front Microbiol. 2017;8:226.

Cunha CS, Veloso CM, Marcondes MI, Mantovani HC, Tomich TR, Pereira LGR, et al. Assessing the impact of rumen microbial communities on methane emissions and production traits in Holstein cows in a tropical climate. Syst Appl Microbiol. 2017;40:492–9.

Ramayo-Caldas Y, Zingaretti L, Popova M, Estellé J, Bernard A, Pons N, et al. Identification of rumen microbial biomarkers linked to methane emission in Holstein dairy cows. J Anim Breed Genet. 2020;137:49–59.

Friedman N, Jami E, Mizrahi I. Compositional and functional dynam-
ics of the bovine rumen microecosystem across different development stages. Environ Microbiol. 2017;19:3365–73.

Poulsen M, Schwab C, Borg Jensen B, Engberg RM, Søergaard N, et al. Methanogenic and bacterial community structure. Front Microbiol. 2017;8:226.

Cunha CS, Veloso CM, Marcondes MI, Mantovani HC, Tomich TR, Pereira LGR, et al. Assessing the impact of rumen microbial communities on methane emissions and production traits in Holstein cows in a tropical climate. Syst Appl Microbiol. 2017;40:492–9.

Ramayo-Caldas Y, Zingaretti L, Popova M, Estellé J, Bernard A, Pons N, et al. Identification of rumen microbial biomarkers linked to methane emission in Holstein dairy cows. J Anim Breed Genet. 2020;137:49–59.

Friedman N, Jami E, Mizrahi I. Compositional and functional dynam-
ics of the bovine rumen microecosystem across different development stages. Environ Microbiol. 2017;19:3365–73.

Poulsen M, Schwab C, Borg Jensen B, Engberg RM, Søergaard N, et al. Methanogenic and bacterial community structure. Front Microbiol. 2017;8:226.

Cunha CS, Veloso CM, Marcondes MI, Mantovani HC, Tomich TR, Pereira LGR, et al. Assessing the impact of rumen microbial communities on methane emissions and production traits in Holstein cows in a tropical climate. Syst Appl Microbiol. 2017;40:492–9.

Ramayo-Caldas Y, Zingaretti L, Popova M, Estellé J, Bernard A, Pons N, et al. Identification of rumen microbial biomarkers linked to methane emission in Holstein dairy cows. J Anim Breed Genet. 2020;137:49–59.

Friedman N, Jami E, Mizrahi I. Compositional and functional dynam-
ics of the bovine rumen microecosystem across different development stages. Environ Microbiol. 2017;19:3365–73.

Poulsen M, Schwab C, Borg Jensen B, Engberg RM, Søergaard N, et al. Methanogenic and bacterial community structure. Front Microbiol. 2017;8:226.

Cunha CS, Veloso CM, Marcondes MI, Mantovani HC, Tomich TR, Pereira LGR, et al. Assessing the impact of rumen microbial communities on methane emissions and production traits in Holstein cows in a tropical climate. Syst Appl Microbiol. 2017;40:492–9.
83. Difford GF, Plchta DR, Lewandahl P, Lassen J, Noij SJ, Højberg O, et al. Host genetics and the rumen microbiome jointly associate with methane emissions in dairy cows. PLoS Genet. 2018;14:e1007580.

84. Min BR, Castleberry L, Allen H, Parker D, Waldrip H, Brauer D, et al. Associative effects of wet distiller's grains plus solubles and tannin-rich peanut skin supplementation on in vitro rumen fermentation, greenhouse gas emissions, and microbial changes. J Anim Sci. 2019;97:4668–81. https://doi.org/10.1093/jas/sky2317.

85. Wang M, Wang R, Xie TY, Janssen PH, Sun XZ, Beauchemin KA, et al. Shifts in rumen fermentation and microbiota are associated with dissolved ruminal hydrogen concentrations in fermenting dairy cows fed different types of carbohydrates. J Nutr. 2016;146:1714–21. https://doi.org/10.3945/jn.116.232462.

86. Wang M, Wang R, Zhang X, Ungerfeld EM, Long D, Mao H, et al. Molecular hydrogen generated by elemental magnesium supplementation alters rumen fermentation and rumen microbiota in goats. Br J Nutr. 2017;118:401–10. https://doi.org/10.1017/S0007114517002161.

87. Latham EA, Anderson RC, Pinchak WE, Nisbet DJ. Insights on alterations to the rumen ecosystem by nitrate and nitrocompounds. Front Microbiol. 2016. https://doi.org/10.3389/fmicb.2016.00228.

88. Van Zijderveld SM, Gerrits WJJ, Apaajathi JA, Newbold JR, Dijkstra J, Leng RA, et al. Nitrate and sulfate: effective alternative hydrogen sinks for mitigation of ruminal methane production in sheep. J Dairy Sci. 2010;93:5856–66.

89. Granja-Salecedo YT, Fernandes RM, De Araujo RC, Kushi LT, Berchelli TT, De Resende FD, et al. Long-term encapsulated nitrate supplementation modulates ruminal microbial diversity and rumen fermentation to reduce methane emission in grazing steers. Front Microbiol. 2019;10:614.

90. Shah AM, Ma J, Wang Z, Hu R, Wang X, Peng Q, et al. Production of hydrogen sulfide by rumen fermentation and its impact on health and productivity of animals. Processes. 2020;8:1169.

91. Fonty G, Joblin K, Chavarot M, Roux R, Naylor G, Michallon F. Establishment and development of ruminal hydrogenotrophs in methanogen-free lambs. Appl Environ Microbiol. 2007;73:6391–403. https://doi.org/10.1128/AEM.00181-07.

92. Chen L, Luo Y, Wang H, Liu S, Shen Y, Wang M. Effects of glucose and starch on rumen fermentation by newly isolated Streptococcus bovis 51 from Saanen Goats. Appl Environ Microbiol. 2016;82:5982–9. https://doi.org/10.1128/AEM.01994-16.

93. Zhang R, Liu J, Jiang L, Mao S. Effect of high-concentrate diets on microbial composition, function, and the VFAs formation process in the rumen of lactating dairy cows. J Anim Feed Sci. 2020;29:114619.

94. Prabhu R, Altman E, Eiteman MA. Lactate and acrylate metabolism by Succiniclasticum ruminis gen. nov., sp. nov., a ruminal hydrogenotroph. J Microbiol. 2012;48:67–76. https://doi.org/10.1007/s12275-011-1141-7.

95. Shabat SKB, Sasson G, Doron-Faigenboim A, Durman T, Yaacoby S, Berg Miller M, et al. Nitrate and sulfate: effective alternative hydrogen sinks for mitigation of ruminal enterobacteria infection, ruminal cellulolytic bacteria populations and cellulase activities of corn stover sludge. J Integr Agr. 2020;19:383–97. https://doi.org/10.1016/S2095-319X(19)62707-3.

96. Jeyanathan J, Martin C, Eugène M, Ferlay A, Popova M, Morgavi DP. Bacterial direct-fed microlabs fail to reduce methane emissions in primiparous lactating dairy cows. J Anim Sci Biotechnol. 2019;10:141. https://doi.org/10.1186/s40104-019-0342-9.

97. pope PB, Smith W, Denman SE, Triinge SG, Barry K, Hugenholtz P, et al. Isolation of Succinivibrioaceae implicated in low methane emissions from Tammar Wallabies. Science (80)-al. 2011;335:646–8. https://doi.org/10.1126/science.1205760.

98. Scottenbach JR, Kristensen NB, Donkin SS, Hammon HM, Penner GB. Gluconeogenesis in dairy cows: the secret of making sweet milk from sour dough. JUBMB Life. 2010;62:869–77.

99. Armstrong DE, Motilova M, Seibertova O, Balasekova L, Scurtova L, Kajper J, et al. Factors influencing rumen fermentation efficiency of hydrogen on formation of propionate. J Dairy Sci. 1998;81:1165–7. https://doi.org/10.3168/jds.S0022-0302(76)84383-4.

100. Pope PB, Smith W, Denman SE, Tringe SG, Barry K, Hugenholtz P, et al. Isolation of Succinivibrioaceae implicated in low methane emissions from Tammar Wallabies. Science (80)-al. 2011;335:646–8. https://doi.org/10.1126/science.1205760.

101. Santry JG, Fasina D, Gomel D, Wang M, et al. Nitrate and sulfate: effective alternative hydrogen sinks for mitigation of ruminal acetate bacteria to reduce methane production in ruminants, a critical review. Front Microbiol. 2010;10:2207.

102. Chen J, Harstad OM, McAllister T, Dorsch P, Holan H. Propionic acid bacteria enhance ruminal fixed feed degradation and reduce methane production in vitro. Acta Agric Scand A Anim Sci. 2020;69:169–75. https://doi.org/10.1080/09006470.2020.1773215.