Hydrogen and formate production and utilisation in the rumen and the human colon

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Abstract
Molecular hydrogen (H₂) and formate (HCOO⁻) are metabolic end products of many primary fermenters in the mammalian gut. Both play a vital role in fermentation where they are electron sinks for individual microbes in an anaerobic environment that lacks external electron acceptors. If H₂ and/or formate accumulate within the gut ecosystem, the ability of primary fermenters to regenerate electron carriers may be inhibited and microbial metabolism and growth disrupted. Consequently, H₂- and/or formate-consuming microbes such as methanogens and homoacetogens play a key role in maintaining the metabolic efficiency of primary fermenters. There is increasing interest in identifying approaches to manipulate mammalian gut environments for the benefit of the host and the environment. As H₂ and formate are important mediators of interspecies interactions, an understanding of their production and utilisation could be a significant entry point for the development of successful interventions. Ruminant methane mitigation approaches are discussed as a model to help understand the fate of H₂ and formate in gut systems.

Keywords: Hydrogen, Formate, Rumen, Colon, Methane, Methanogens, Homoacetogens, Mitigation

Introduction
Organoheterotrophic anaerobic microorganisms derive their energy by breaking down biomass and fermenting the monomers and oligomers that are released. In anoxic environments this fermentation is typically catalysed by syntrophic interactions between microorganisms. The anaerobic environments in mammalian gut ecosystems are microbial habitats that differ from non-gut systems in that they have relatively short digesta residence times with turnover once or twice per day, compared to anaerobic bioreactors (>14–20 days) and sediments (years and decades). The faster turnover in gut systems results in incomplete fermentation with the generation of mainly volatile fatty acids (VFAs), methane (CH₄), and carbon dioxide (CO₂). There is also a need for relatively fast ATP generation so that microbes can grow fast enough so they are not washed out of the system. The VFAs are absorbed from the gut and serve as energy substrates and metabolites for the host animal while CH₄ and CO₂ are emitted as gaseous wastes [1]. In the foregut (rumen) of ruminants and the human large intestine CH₄ and CO₂ account for around 10 and 17% of the fermentable carbon respectively, with the remainder found as VFAs [2, 3]. In contrast, in biodigesters and sediments where VFAs are also metabolised, organic material is completely degraded to CH₄ and CO₂ [1].

In 1981 Meyer Wolin [3] reviewed what was then known about fermentation in the rumen and the human large intestine. This review highlighted the role of H₂ and formate production and utilisation in these environments, and the importance of interactions between...
different microbes in interspecies H2 transfer. It also discussed approaches to reduce CH4 production in order to improve animal performance and feed utilisation. Since that time our knowledge of gut environments has rapidly increased. While the most significant development has been determining the diversity of microbial species in these gut environments, there have been advances in understanding the overall metabolism of the microbes and the enzymes encoded in their genomes. Besides interspecies H2 transfer, interspecies formate transfer has been realized to be of importance in mediating electron flow [4]. Also, it has been shown that the gut environment is not homogeneous but consists of planktonic and aggregated cell fractions (biofilms) associated with particulate matter and gut surfaces that have different passage rates and in which interspecies electron flow is significantly different [5, 6]. In recent years emphasis has been placed on research to understand the contribution of the gut microbiome to human health and to mitigate the environmental effects of ruminant CH4 emissions. However, there is still much to learn about how gut microbes interact with each other, with dietary components and with their mammalian host. In this review we examine mutualistic fermentative digestion in the ruminant anterior forestomach (rumen-reticulum or rumen) and the distal human colon emphasising the role that H2 and formate production and use play in their metabolisms. We also discuss progress in interventions that could potentially redirect H2 and formate metabolism and impact ruminant methane production.

**Differences between the two gut environments**

The ruminant rumen is a large, pre-gastric fermentation organ in which mutualistic microbial fermentation takes place prior to gastric digestion. In contrast, the human colon is a much smaller, post-gastric fermentation chamber. Important features of these two gut fermentation systems are listed in Table 1. For studies of the human gut most samples are of faecal material or are obtained via other non-invasive techniques, whereas for ruminants collection of digesta directly from the rumen of cannulated animals or by oral intubation is well established [7, 8]. Furthermore, in ruminants the composition of the diet and the administration of feed additives can be precisely controlled making scientific experimentation

| Characteristic                  | Rumen                                      | Human colon                                      |
|--------------------------------|--------------------------------------------|-------------------------------------------------|
| Mode of digestion              | Pregastric—foregut fermentor              | Postgastric—hindgut fermentor                   |
|                                | Continuously stirred/mixed tank reactor   | Plug flow tubular system                        |
| Diet                           | Evolved for efficient fibre degradation and utilisation. Major metabolizable energy supply (~ 70%) of the host energy requirements | Adapted to hydrolysing and fermenting undigested dietary residues and host endogenous secretions. Minor contribution to host energy requirements |
|                                | Breakdown dietary protein and non-protein nitrogen for synthesis of microbial protein | Post absorptive compartment                      |
|                                | Synthesis of B vitamins                    | Post absorptive compartment                      |
|                                | Rumen system evolved for detoxification/biotransformation of phytotoxins and mycotoxins | Host system evolved for transport and excretion of toxic/xenobiotic compounds |
| Blood glucose                  | Low—rely on gluconeogenesis to generate glucose precursors | High                                           |
| Microbiology                   | Anaerobic bacteria and methanogenic archaea | Anaerobic bacteria and methanogenic archaea     |
|                                | Ciliate rumen protozoa                     | Flagellate protozoa?                            |
|                                | Anaerobic rumen fungi                      |                                                |
|                                | Bacteriophage                              | Bacteriophage                                   |
| VFA/SCFA                       | Acetate, propionate and butyrate are the predominant volatile fatty acids | Similar molar proportions of the three main volatile fatty acids |
|                                | Branched chain VFAs                       | Similar                                         |
|                                | Lactate → Propionate                       | Similar turnover                                |
|                                | Succinate → Propionate                     |                                               |
| Gas composition                | CO2 65%; CH4 27%; N2 7%; O2 0.6%; H2 0.2% | CO2 10%; CH4 14%; N2 65%; O2 2.3%; H2 3%       |
|                                | CO2 produced from fermentation and HCO3− in saliva. N2 and O2 is ingested with feed and diffuses through the rumen wall. Partial pressure of H2 maintained at a very low level. CH4 emission amounts to 2–12% of gross energy | NO2 and O2 ingested, with CO2, H2 and CH4 resulting from colonic fermentation. Less CO2 and CH4 than the rumen but with higher H2 concentrations |
|                                | Gas elimination                            | Flatus and reabsorption and removal by lungs    |
|                                | Mainly eructation                          |                                               |
stronger and more rigorous than the investigation of the human colon.

Both environments harbour dense and diverse populations of microorganisms that form closely integrated ecological units with their hosts. The rumen is inhabited by a diverse microbial community comprised of anaerobic bacteria, methanogenic archaea, ciliate protozoa, anaerobic phycocyanin fungi and bacteriophage and is known to be highly adaptable metabolically to deal with changes in diet. One main difference of the rumen compared to the human colon is the presence of a large eukaryal population of ciliate protozoa that accounts for as much as 50% of the microbial biomass. Under normal healthy conditions, the human colon mainly anaerobic bacteria, methanogenic archaea, and bacteriophage. In both systems, the microbial inhabitants play vital roles in the nutritional, physiological, immunological, protective, and developmental functions of their respective hosts, but the forces that control and shape the composition and activities of these microbial communities remain poorly understood.

A major difference between the rumen and the colon is that in the rumen the microbes initiate feed degradation, while in the colon the host digestive processes act on the feed first. The diet of farmed ruminants is largely composed of fibre (cellulose, hemicelluloses, and pectin) and starch in varying proportions depending on the production system with a relatively constant daily intake. The human diet is highly variable and the fermentation substrates which reach the colon include undigested dietary polysaccharides such as fibre, resistant starch, and oligosaccharides that escape digestion in the upper tract. Host-secreted mucin glycans are also an important substrate for human gut microbes [9]. Rumen microbes are not thought to use host glycans, but the presence of host glycan-degrading enzymes in some rumen Prevotella spp. [10] suggests they may be able to use salivary glycoproteins.

Acetic, propionic, and butyric acids are the major VFA products of fermentation in both the rumen and human colon. It is well established that rumen VFAs are absorbed and contribute about 70% of the animals metabolizable energy requirement [11]. VFAs are also absorbed from the human large intestine and contribute to energy requirements of the host albeit at a much lower level (~ 10%, [11]). An important difference lies in the production of gaseous products. Intestinal gases of humans [12] have a lower percentage of CO₂ and CH₄ and a greater percentage of H₂ than gases found in the rumen. CH₄ emission is universal in rumen fermentation, whereas the proportion of humans identified as CH₄ emitters varies [13, 14], with 20% of Western populations identified as high emitters [15]. Moreover, H₂ is rarely a final product of rumen fermentation but is always a product of large intestinal fermentation in humans and significant amounts of residual H₂ that is not used by microbes are excreted via expiration or flatus.

**Hydrogen and formate metabolism in the rumen**

H₂ is primarily produced during microbial fermentation by hydrogenases. These enzymes catalyse the reoxidation of cofactors reduced during carbohydrate fermentation [16] and dispose of the derived electrons by reducing protons to produce H₂. In the rumen most of the H₂ produced is used by methanogenic archaea to reduce CO₂ (hydrogenotrophic methanogenesis) or methyl compounds (methylo trophic methanogenesis) to CH₃ via a process known as interspecies hydrogen transfer [17]. H₂ is maintained at sufficiently low concentrations through methanogenesis for fermentation to remain thermodynamically favourable [16, 18].

A range of rumen microbes belonging to several different phyla have been shown to produce H₂, with 65% of cultured rumen bacterial and archaeal genomes [10] encoding enzymes that catalyse H₂ production or consumption [19]. Metagenome assembled genomes (MAGs) from different gastrointestinal tract regions of seven ruminant species [20] generated similar results. A total of 6,152 [NiFe]-, [FeFe]-, and Fe-hydrogenase-containing MAGs were detected, 3003 of which encoded enzymes for fermentative H₂ production (72.7% from the Firmicutes), while 95 MAGs encoded H₂-uptake hydrogenases and the methyl-CoM reductases related to hydrogenotrophic methanogenesis (mainly from Methanobrevibacter).

Flavin-based electron bifurcation is an electron pair-splitting mechanism that enables the coupling of energy-producing redox reactions with energy-consuming electron transfer reactions [21] and is likely to be particularly important for fermentation in the anaerobic gut environment. Metatranscriptomic analysis using data from sheep that differed in their methane yield [22] showed that electron-bifurcating [FeFe]-hydrogenases were key mediators of ruminal H₂ production [19]. Hydrogenases from carbohydrate-fermenting Clostridia (Ruminococcus, Christensenellaceae R-7 group) accounted for half of all hydrogenase transcripts, suggesting that these organisms generate much of the H₂ used by the hydrogenotrophic Methanobrevibacter species. Co-culturing experiments showed that the hydrogenogenic cellulose fermenter Ruminococcus albus expressed its electron-bifurcating hydrogenase and suppressed its ferredoxin-only hydrogenase when grown with the hydrogenotrophic fumarate reducer Wolinella succinogenes [19, 23].
Rumen methanogens also participate in symbiotic relationships with protozoa that produce large quantities of H$_2$ via their hydrogenosomes [24]. In return, the protozoa benefit from H$_2$ removal as high H$_2$ partial pressure is inhibitory to their metabolism. Meta-analysis of protozoa defaunation studies concluded that elimination of ciliate protozoa reduced CH$_4$ production by up to 11% [25]. A similar relationship exists between methanogens and anaerobic rumen fungi which also contain hydrogenosomes [26].

Metatranscriptomic studies [19] showed that, while enzymes mediating fermentative H$_2$ production were expressed at similar levels, methanogenesis-related transcripts predominated in high methane yield sheep, while alternative H$_2$ uptake pathways were significantly upregulated in low methane yield sheep. These other H$_2$ uptake pathways could potentially limit CH$_4$ production by redirecting H$_2$ uptake away from methanogenesis towards homoacetogenesis (Blautia, Eubacterium), fumarate and nitrite reduction (Selenomonas, Wolinella), and sulfate reduction (Desalfovibrio). Homoacetogens produce acetate from H$_2$ and CO$_2$ and are known to occur in the rumen, but their abundance is generally lower than hydrogenotrophic methanogens [27, 28]. It is likely that methanogens outcompete homoacetogens at the low H$_2$ concentrations in the rumen [29]. Nitrate and sulfate reduction are thermodynamically more favourable than methanogenesis and homoacetogenesis [16], and nitrate and sulfate-reducing bacteria occur naturally in the rumen. Their population densities increase as the concentration of their respective electron acceptors in the ruminant diet increases. However, nitrate and sulfate concentrations in ruminant diets are usually very low so these processes would be substrate limited and their end products can be toxic at high concentrations [30].

Much less is known about formate concentrations in the rumen or the significance of formate as an electron carrier between species [31]. At relatively high redox potentials (low H$_2$ and formate concentrations), formate is thermodynamically and kinetically a more favourable interspecies electron carrier than H$_2$. This is of importance mainly for planktonic microbes, where the distances for electron transfer between organisms are greater than between those growing in biofilms and aggregates [5]. Many rumen microbes contain formate dehydrogenase genes, but their expression under different conditions has been much less studied compared to hydrogenase genes.

**Hydrogen and formate metabolism in the human colon**

Less is understood about the H$_2$ and formate economy of the human gastrointestinal tract. H$_2$ is formed in large volumes in the colon as an end product of polymeric carbohydrate fermentation, for example by members of the Firmicutes and Bacteroidetes. Two major pathways for disposal of H$_2$ are known, methanogenesis, and homoacetogenesis, with homoacetogenesis being the predominant pathway [32, 33]. When sulfate is available, dissimilatory sulfate reduction is carried out by sulfate-reducing bacteria but in its absence these bacteria thrive by producing H$_2$ rather than oxidizing it. Most methanogens require H$_2$ or formate for growth whereas homoacetogens are metabolically versatile and can also utilise a wide range of organic substrates. The dominance of one pathway over another appears to vary among individuals and to be inherited; there are individuals in which most of the H$_2$ generated is converted to CH$_4$ and in others H$_2$ can accumulate to high concentrations [12]. Spatial and temporal variations in the chemical composition of the digesta in the human colon could result in specific microhabitats that support different H$_2$ and/or formate metabolizing microbes. Homoacetogenesis is energetically less favourable than methanogenesis but may offer greater benefits to the host as CO$_2$ and H$_2$ are converted to acetate for use as an energy source, with no evolution of gas. Microbes using these two pathways represent key species in the human gut microbiome [34].

Although the formation and use of H$_2$ and/or formate in some of the colonic microbes are well studied in pure culture, the hydrogenases and formate dehydrogenases responsible have not been investigated and our understanding of their roles at different stages of the fermentation process is poor. Only one study has surveyed the genomic and metagenomic distribution of hydrogenase-encoding genes in the human colon to infer the dominant mechanisms of H$_2$ cycling [35]. Most microbial species from the Human Microbiome Project Gastrointestinal Tract reference genome database encoded the genetic capacity to form or use H$_2$. A wide variety of anaerobically adapted hydrogenases were present, with [FeFe]-hydrogenases predominant. Metagenomic analysis of stool samples from 20 healthy humans indicated that the hydrogenase gene content of all samples was overwhelmingly dominated by fermentative and electron-bifurcating [FeFe]-hydrogenases from members of the Bacteroidetes and clostridial members of the Firmicutes. This study concluded that H$_2$ metabolism in the human colon is driven by fermentative H$_2$ production and interspecies H$_2$ transfer using electron-bifurcation rather than respiration as the dominant mechanism of H$_2$ reoxidation. However, microenvironmental niches in this mucosal ecosystem have not been fully studied and both the cellular and genetic bases for H$_2$ and formate cycling in the human colon require further exploration.

Bacteria belonging to the family Christensenellaceae are highly heritable members of the human gut microbiome.
that show strong correlations with host health [36]. Analysis of human gut metagenomes have identified positive associations between Christensenella spp. and Methanobrevibacter smithii. In co-cultures these organisms grow together in dense flocs, and H₂ generated by Christensenella spp. supports CH₄ production by M. smithii [37]. This interaction shifts fermentation end products toward more acetate and less butyrate, potentially affecting the physiology of the human host. Interaction between M. smithii and members of the Christensenellaceae R-7 group has also been highlighted in analysis of the microbiomes of 30 subjects identified as high or low CH₄ emitters [38]. High emitters were characterised by a 1000-fold increase in M. smithii which co-occurred with bacteria from a core group of keystone species from the Christensenellaceae and Ruminococcaceae families. Cross feeding between H₂ and formate-producing bacteria and the human gut homoacetogen Blautia hydrogenotrophica has also been demonstrated in co-cultures [38–40].

Redirecting hydrogen metabolism and ruminant methane mitigation

While there is interest in trying to determine if a mechanistic/causative relationship exists between the presence of methanogens and human gut health [15, 41], there has been increasing urgency in developing approaches that can practically mitigate CH₄ from ruminant animals [42]. Globally, CH₄ from enteric fermentation in ruminant livestock is a major source of agricultural greenhouse gases [43, 44]. Ideally, any developed mitigation approach should induce a co-benefit for the animal, for example enhanced production or health. Co-benefits can help drive practical adoption of the technology on farm. While research into reducing ruminant CH₄ emissions has been in progress for many years [45, 46] and promising mitigation approaches are being developed [47], emergence of co-benefits from these approaches is not being observed consistently. This is contrary to the frequently stated hypothesis that ruminal CH₄ production represents a loss of energy, from 2 to 12% of gross energy intake [48], which could in principle otherwise be available for animal growth or milk production. Historically, it has also been hypothesized that H₂ accumulation resulting from the inhibition of methanogenesis will impair fibre digestion and fermentation [18]. It is becoming clear that a lack of understanding of H₂ and formate metabolism and how it can be manipulated is a barrier that needs to be overcome in order to support the development of CH₄ mitigation approaches with co-benefits. Emerging CH₄ mitigation strategies in ruminants are now available and can provide model systems to help advance our knowledge in this area. Four promising areas are discussed below.

Animal selection and breeding

Animals vary in their methane production and breeding low methane emitting animals is one mitigation approach. Significant progress has been made with sheep where studies have found animals that vary naturally in the amount of CH₄ they produce. The heritability of this trait has enabled the breeding of low-CH₄ emitting sheep [49, 50], and CH₄ emissions from selected, divergent lines differ on average by 10–12%. Physiological characteristics such as a reduction in the rumen retention time of feed particles [51] and reduced rumen volume [52] are factors likely to contribute to the low CH₄ emissions. There are also differences observed in their rumen microbial communities [53] and expression of microbial genes involved in the production of CH₄ are reduced in low-CH₄ sheep [22]. Kamke et al. [54] proposed that the rumen microbiome in low-CH₄ animals supported heterofermentative growth leading to lactate production, with the lactate subsequently metabolised mainly to butyrate. Greening et al. [19] offered an alternative interpretation with H₂ uptake through non-methanogenic pathways accounting for the differences observed.

Competing terminal electron acceptors or alternative H₂ users

Several alternative electron acceptors have been added to ruminant diets in attempts to alter the rumen fermentation and reduce CH₄ production. Nitrate is the most studied compound [46] and is reduced via nitrite to ammonia, reducing the availability of H₂ for CH₄ synthesis. Sulfate reduction will also compete for electrons and H₂ and may lower CH₄ production [30]. Stimulating the activity of acetogens through the inhibition of methanogens has been proposed as a strategy for ruminant CH₄ mitigation [55], but it is unclear whether resident rumen homoacetogens could fulfil the H₂ disposal role or whether non-resident homoacetogens would need to be inoculated into the rumen [56]. Studies to date where methanogenesis has been inhibited with an effective methane inhibitor, such as 3-nitrooxypropanol, have not demonstrated increased homoacetogenesis.

In the gut of macropod marsupials, which consume a diet similar to ruminants, CO₂ is mainly reduced to acetate rather than to CH₄ as a means of electron disposal [28]. The reason for the preference of this alternative pathway in macropods is unknown, but it has been argued that their tubiform forestomach lacks the mechanisms to remove gaseous products of fermentation (e.g., eructation in the rumen and flatus in the lower bowel) and their immune secretions suppress the microbes responsible for releasing H₂ or CH₄ to prevent gas pressure build-up that would threaten gut integrity [57].
Methanogen inhibiting technologies

Several different approaches have been used to specifically target methanogens in the rumen. These include feed additives such as 3-nitrooxypropanol [58], halogenated compounds [59], and certain seaweeds [60] as well as work to develop anti-methanogen vaccines [61]. Studies to date with the available technologies suggest that when CH$_4$ production is inhibited, we do not observe a sufficient increase in rumen H$_2$ emissions to account for the reducing equivalents that are not captured in CH$_4$. It is assumed that the electrons are being diverted to other fermentation products, such as acetate, propionate, butyrate, and microbial biomass, but the balance of this redirection of electrons is not well understood. The use of methanogen inhibitors in combination with microbes that could potentially redirect H$_2$ to other products has yet to be explored.

An alternative approach is to target the microbes that produce the substrates for methanogenesis. Although recent work has begun to identify the bacteria most likely to produce H$_2$ [19] and methyl compounds [62] used as substrates for methanogenesis in the rumen, significant knowledge gaps remain. At this point it is unknown if a reduction in the production of substrates for methanogenesis would impede overall fermentation.

Diet

Although methane emissions arising from an individual animal are primarily driven by the quantity of feed eaten [63], the chemical composition of feeds can also influence emissions. Consequently, the nature of the feed consumed may select for microbial populations with different fermentation pathways that yield less H$_2$ and therefore less CH$_4$. For example, concentrate-based diets are associated with lower CH$_4$ yield (g/kg DMI; [48]) because fermentation of starch in concentrate results in more propionate and butyrate being produced and less CH$_4$. Fermentation products [64] and microbial composition [65] may be influenced by the oxidation state of the carbon substrates especially those with higher levels of the more reduced sugar alcohols or more oxidised sugar acids. Brassica forages have also been shown to result in lower CH$_4$ yields in lambs than perennial ryegrass [66]. The reason for this is not understood but an altered rumen microbiota or the presence of bioactive glucosinolates in brassicas have been suggested as possible causes [67]. Generally, however, it takes large changes in diet to bring about significant changes in enteric methane emissions in ruminants.

Conclusion

Here we have contrasted the rumen and human colon environments and H$_2$ and formate have emerged as key metabolites involved in cross-feeding between members of the microbiota in both gut ecosystems with important roles in shaping the syntrophic networks that operate in these environments. The role of H$_2$ and formate production as electron sinks for individual microbes and their transfer of electrons to homoacetogenic bacteria and methanogenic archaea are key functions to ensure ongoing polysaccharide degradation and energy generation for both ruminants and humans. Although the anaerobic bacteria and archaea are broadly similar in each environment, H$_2$ produced in the rumen is consumed predominantly by incorporation into CH$_4$, whereas in the human colon significant H$_2$ emissions escape the system. Determining what controls these differences will be important in understanding the impact of H$_2$ and formate turnover on human gut metabolism and in reducing the environmental impact of ruminant CH$_4$. Currently, the availability of emerging CH$_4$ mitigation approaches for ruminant animals makes the rumen an ideal gut system to study the production and utilisation of hydrogen and formate in gut systems and generate knowledge applicable to both systems.

Our present knowledge of the mammalian gut H$_2$ and formate economy is incomplete and an improved understanding of the active groups of microbes involved in H$_2$, and formate metabolism is required. Cultures and genome sequences of model hydrogenotrophs, such as *Methanobrevibacter* and *Blautia*, are available and have been used to demonstrate interspecies H$_2$ and formate transfer in co-culture. However, our knowledge of which organisms produce the bulk of the H$_2$ and formate in gut environments is limited. The metagenome- and metatranscriptome-based studies have highlighted the diversity of gut microbes encoding the signature genes for hydrogenotrophy and have emphasized the need for more exact information about the function of these genes in the gut environment. There also remains a need to bring a greater proportion of representatives of the currently uncultured microorganisms into cultivation together with additional host-associated homoacetogen and methanogen strains. This should be accompanied by studies of their physiology, metabolism, and interactions with other gut anaerobes to provide a body of knowledge beyond what can be inferred from genome and metagenome sequence data. With the increased availability of such pure cultures and their corresponding genome sequences it will prove possible to construct metabolically interacting microbial consortia so that the contributions of different microbes to overall community function can be ascertained.
Abbreviations

\( \text{H}_2 \): Hydrogen; \( \text{HCOO}^- \): Formate; VFAs: Volatile fatty acids; \( \text{CH}_4 \): Methane; \( \text{CO}_2 \): Carbon dioxide; MAGs: Metagenome assembled genomes; DMI: Dry matter intake.

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