The rumen microbiome: a crucial consideration when optimising milk and meat production and nitrogen utilisation efficiency

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ABSTRACT
Methane is generated in the foregut of all ruminant animals by the microorganisms present. Dietary manipulation is regarded as the most effective and most convenient way to reduce methane emissions (and in turn energy loss in the animal) and increase nitrogen utilization efficiency. This review examines the impact of diet on bovine rumen function and outlines what is known about the rumen microbiome. Our understanding of this area has increased significantly in recent years due to the application of omics technologies to determine microbial composition and functionality patterns in the rumen. This information can be combined with data on nutrition, rumen physiology, nitrogen excretion and/or methane emission to provide comprehensive insights into the relationship between rumen microbial activity, nitrogen utilisation efficiency and methane emission, with an ultimate view to the development of new and improved intervention strategies.

Introduction

Global dairy sector
The global agri-food industry employs over 1 billion people, representing 1 in 3 of all workers. In 2013, the largest producers of dairy milk were the US, EU and India, with global production levels at 617 billion litres of milk. According to, the EU is expected to become one of the world’s leading exporters of milk, with sales anticipated to rise to 19.2 million tonnes of milk equivalent. Since 2005, milk supply and demand has increased by 26%, at 2.4% per annum, respectively, with a further growth of 25% expected over the next 10 years. According to the FAO, the aggregate meat consumption of the industrial countries grew at 1.3% per annum in the last 10 years. With the global population set to increase to 9.1 billion by 2050, food production will need to increase by 70% to meet demands for adequate nutrition.

However, with an increase in milk and meat production comes an increase in greenhouse gases (GHG). On a global scale, it has been estimated that livestock production contributes between 9% and 11% of total anthropogenic GHG emissions with approximately 44% of livestock emissions in the form of methane (CH₄).

Measuring and mitigation of methane emissions from livestock is increasing in significance. There are currently a number of methane mitigation strategies being investigated to reduce emissions from ruminants, including better feeds and feeding systems i.e. dietary manipulation of the rumen microbiome, improved genetics and overall health of animals (potentially leading to reduced herd sizes while maintaining milk yields), use of supplements that reduce methane emissions and immunisation against methanogens.

The bovine rumen microbiome
The rumen can be viewed as an anaerobic and methanogenic fermentation chamber that contains microorganisms that have the ability to utilise, and increase the productivity of, cellulolytic feeds (i.e. straw, hay, silage and grass). There are considerable benefits associated with understanding rumen function, as rumen dynamics are almost solely...
responsible for providing nutrients to the host animal. Figure 1 shows the gastrointestinal tract of the bovine animal, where the rumen and its microbiota play a particularly important role in the degradation of feedstuffs. As a result of fermenting feedstuffs, carbon dioxide (CO\textsubscript{2}) and hydrogen (H\textsubscript{2}), which are the main electron acceptors and donors of the ecosystem, are produced in the rumen. Recent developments relating to compounds used to shift fermentation towards more efficient microbial pathways and targeting of specific microorganisms will be discussed later in this review. The rumen microbiome, i.e., the community of microorganisms that inhabits the rumen, is characterised by its high population density, extensive diversity (encompassing bacteria, archaea, protozoa and fungi) and complexity of interactions. The continuous fermentation carried out by these microorganisms leads to ingested compounds being broken down into their subcomponents. There are three intersecting micro-environments found in the rumen that contain these microbes; the liquid phase making up 25% of the microbial mass, the solid phase making up 70% of the microbial mass, and the rumen epithelial cells and protozoa, containing 5% of the microbial mass. A nutritionally balanced diet is important as it provides an environment that maximises the growth and activity of these microbes. A rumen microorganism is anaerobic or facultatively anaerobic, and produces end products that are either utilised directly by the host or by other microorganisms as energy. The pH of the rumen is kept relatively constant, typically 6–7, but may vary depending on diet. Such variations can result in a change in the microbial populations, and the levels of volatile fatty acids (VFAs) produced by them. These fatty acids are of interest as some, such as propionate and butyrate, can be absorbed across the gut wall to serve as an energy source for the ruminant. Buffering of the rumen to maintain a relatively constant pH is facilitated by the large quantity of saliva produced by the ruminant, which is high in sodium and potassium bicarbonate and urea. The saliva is swallowed into the rumen and then absorbed through the rumen walls. Further buffering is provided by ammonia produced during fermentation, which can then be used for microbial growth in the rumen.

Ruminants, through the action of their microbiota, can utilize components that the human body cannot break down, namely lignocellulose. Lignocellulose is the most abundant carbon polymer on the planet, with the rumen having a central role in releasing this vast energy store. The rumen ultimately uses lignocellulose to make products (i.e. milk and meat) that are then available to humans to consume as a nutrient dense food source. The interaction between the host and microbes in the rumen is synergistic, in that the host provides heat, moisture and food, while the microorganisms produce protein and by-products of digestion, such as the aforementioned VFAs, for use by the host. The degradation of microbes by the host has also been described in literature, with microbes utilised for their protein, lipid and starch content. The complex rumen ecosystem consists of bacteria, archaea, ciliate protozoa, fungi, bacteriophage and viruses, which will be discussed below.

**Rumen bacteria**

Bacteria are the most abundant microbes in the foregut of ruminant animals, with approximately $10^{10} - 10^{11}$ cells/ml and over 200 species. The composition of the bacteria found in the rumen is dictated by a number of factors including preference for certain substrates (i.e. diet), energy requirements, and resistance to certain metabolic end-products that may be toxic to some species. In a study by Hungate, in which rumen microbial populations were studied using a culture-
based approach, it was determined that rumen bacterial composition is mainly Gram negative when animals are being fed high forage diets, with more Gram positive bacteria, such as *Lactobacillus*, present in animals fed high grain diets, with ruminal pH levels dropping after the consumption of easily fermented carbohydrates.  

The use of molecular techniques has become critical for the analysis of rumen microbiology. Such techniques can be used to determine the composition of the population present (using, for example, the conserved 16S rRNA gene to determine bacterial composition), predict their functionalities or enumerate targeted microbes within a complex ecosystem without the need for culturing. This is important, as it is estimated that only 20% of the rumen microbiota can be cultured using standard techniques. Since sequencing of the 16S rRNA gene was first used to study rumen microbial ecosystems, extensive coverage of low abundance species has enabled the analysis of rare microbial communities. Sequencing also showed that *Prevotella*, *Butyrivibrio* and *Ruminococcus* were the most dominant bacteria in the rumen, and that community structure is affected by changes in the diet of the host. In particular, it has been shown that complexity in the diet favours increased microbial diversity.

Due to the high forage diets of ruminants, particularly those in receipt of grass-based diets, cellulose digesters are an important part of supplying the animal with vital nutrients. These bacteria degrade cellulose and hemicellulose, the main components of plant fibre. The ability to degrade cellulose is strongly dependent on the type of forage, crop maturity and the accessibility of the cellulolytic bacterial communities. The matrix of plant fibre is complex, composed of β-1, 4 linked glucose residues for cellulose and β-1, 4 linked xylose for hemicellulose, requiring the coordination of a number of hydrolytic enzymes in order to break it down. Although there are many cellulose degrading bacteria, *Fibrobacter succinogenes* and *Ruminococcus albus* are the most desirable cellulose degraders. Their ability to digest cellulose is much higher than that of other cellulolytic bacteria, which is probably due to the fact that they possess a number of genes encoding enzymes involved in fibre degradation. Fermentation end products of cellulolytic bacteria include acetate, butyrate, propionate and CO₂. Hydrogen, ethanol, succinic acid, formic acid and lactic acid are also formed but are quickly used by other bacteria.
percentages. Similar results were found by Bainbridge et al. (2016), where moderate correlations were found between milk yield, protein percentage, fat yield and bacterial communities. This alone indicates the role of the rumen microbiome in shaping host physiological parameters.  

**Methanogenic archaea**

Archaea, in general, have a broad spectrum of unusual and distinctive metabolisms, enabling them to survive in a variety of different environments. Rumen archaea are strictly anaerobic and are the only known microorganisms present in the rumen capable of producing methane. Such archaea are referred to as methanogens. Archaea are found in the rumen in the range of $10^6$ to $10^8$ cells per ml, accounting for less than 4% of the microbial community. Archaea are found at the bottom of the trophic chain due to their need to use the end products of fermentation as substrates. The domain Archaea is broken into two different kingdoms; Euryarchaeota, consisting of methanogens and extreme halophiles, and Crenarchaeota, consisting of hyperthermophiles and nonthermophiles. Methanogens found in the kingdom Euryarchaeota require a very low redox potential and are among the strictest anaerobes known. According to meta-analysis of global data, 90% of rumen methanogens belong to the following genera: *Methanobrevibacter* (63.2% of methanogen population), *Methanobrevibacter* (7.7% of methanogen population) *Methanosphaera* (9.8%), “Rumen Cluster C”, now referred to as *Thermoplasmata* (7.4%) and *Methanobacterium* (1.2%). Most methanogens remove hydrogen gas by reducing CO$_2$ with hydrogen gas to form methane. In contrast, *Methanosphaera* *stadtmannae* only produces methane through the reduction of methanol with H$_2$, having one of the strictest energy metabolisms of all methanogenic archaea. Producing methane keeps hydrogen concentrations in the rumen low, allowing methanogens to promote the growth of other species, and enabling a more efficient fermentation. However, methane produced in the rumen is eructated, leading to atmospheric pollution. Efforts to mitigate rumen methane emissions include vaccines (targeting rumen methanogens through the generation of antibodies to selected methanogen antigens that enter via saliva, binding to targets on the methanogens), small-molecule inhibitors (targets enzymes essential for the growth of methanogens), additives and breeding approaches. In a study carried out by Goopy et al., it was found that sheep that emitted low methane levels had a smaller rumen in comparison to high methane-emitting sheep. There was no difference in dry matter intake or digestibility between the two groups. The study also found that low methane-emitting animals had a shorter mean retention time for both solid and liquid phase. This may be the basis for breeding animals with a smaller rumen size to reduce methane emissions. However, dietary manipulation is regarded the most effective and straightforward method of lowering rumen methane emissions, as selective breeding is slow and selection of specific traits may affect favourable variants.

It should also be noted that methane production by archaea represents an energy loss of about 2 – 12% of gross energy intake, meaning this energy is no longer available for animal growth, lactation, maintenance or pregnancy. Manipulating the diet of ruminants to reduce the number of methanogens would therefore both help reduce the negative impact on the environment, and also improve the efficiency of livestock production. Methanogenesis is the only mechanism of ATP synthesis available to methanogenic archaea. The methyl coenzyme A reductase (McRA, encoded by mCRA) catalyses the final step in methanogenesis, which means that this enzyme and the genes associated with it may be good markers for the presence of methanogens. Furthermore, anti-methanogens, such as bromochloromethane, can be used to inhibit methane production. Bromochloromethane reacts with reduced vitamin B$_{12}$ and inhibits the cobamide-dependant methyl transferase step, which is responsible for the synthesis of McRA. Targeting this step may break the pathway and inhibit production. A recent study, carried out by Kinley et al., investigated the use of an invasive species of macroalgae as a means of reducing methane emissions *in vitro*. The results showed a decrease in methane production as a result of a secondary metabolite found in the macroalgae *Asparagopsis taxiformis*, bromoform, which is similar...
to bromochloromethane in its ability to decrease methanogenic activity.

**Ciliate protozoa**

Ciliate protozoa are found in the range of $10^4$–$10^6$ cells/ml in rumen fluid and are responsible for 30 to 40% of overall fibre digestion.\(^{16}\) They are also relatively active in lipid hydrolysis and can produce hydrogen via their hydrosomes.\(^{45}\) The *Entodinium* genus is the most dominant protozoan in high grain diets. This genus rapidly degrades starch, engulfing it and converting it to an iodo-philic storage polymer.\(^{16}\) Degradation occurs through a combination of debranching, amylase and glucosidase enzymes. More research may be needed in order to determine their immediate role in methanogenesis and therefore create a better understanding of the value of manipulating this population as a means of reducing methane emissions in ruminants.

Defaunation is a mechanism used to prevent the growth of, or remove, protozoa from the rumen. This can be carried out by several different means such as through the isolation of calves after birth, chemical defaunation through the use of copper sulphate, calcium peroxide, alcohol ethoxylate, coconut oil, linseed oil or soya oil hydrolysate.\(^{46}\) Another method of defaunation, reviewed by Hook et al.\(^{32}\) involves removal of rumen contents from the host, an invasive method whereby the mucosa is carefully washed and the rumen contents are treated by either heating or freezing to eliminate protozoa. The rumen contents are then returned to the rumen. This method has been used to remove protozoa from the rumen to investigate the role they play in rumen function.\(^{32,47}\) suggested that protozoa-associated methanogens (methanogens sequestered within rumen protozoa) which account for approximately 37% of methanogenesis in ruminants,\(^{48}\) may be the most active methanogen communities in the rumen, and as defaunation would also eliminate them, this would account for the decrease in methane production. Conversely, in a study carried out by Morgavi et al.,\(^{8}\) it was concluded that there was no direct correlation between the number of methanogens and amount of methane produced, but there was a correlation between the number of protozoa and methane production (Wallace et al.).\(^{42}\) showed that methane emissions increased as protozoal abundance increased and stated that protozoa showed the strongest link with methane emissions (in comparison to other domains) after qPCR using DNA extracted from rumen contents. Despite the conflicting reports on their roles in methanogenesis, it is agreed that, ultimately, the presence of protozoa can probably have both beneficial and negative effects on the rumen microbiome,\(^{49}\) as protozoa engulf bacteria and other smaller microbes and particles in the rumen, but also larger molecules including proteins and carbohydrates.\(^{50}\) They actively ingest the bacteria as a source of protein and also act as a stabilising factor for fermentation end products.\(^{11}\)

**Amoeba**

Amoebae can represent an important reservoir for bacteria in the environment, but their role in the rumen is unclear. In the vegetative cycle (multiplication by binary fission), amoeba, similar to ciliate protozoa, survive by ingesting bacteria through phagocytosis.\(^{51}\) While further research is necessary in order to ascertain the role of amoeba in the rumen, it is known that some bacteria can survive phagocytosis by protozoa and live as endosymbionts.\(^{52}\) For instance, *Campylobacter jejuni* has been shown to invade *Acanthamoeba polyphaga* and can replicate in vacuoles.\(^{53}\) *C. jejuni* and *C. fetus* can have large effects on cow fertility, immunity and overall health.\(^{54}\) Amoeba therefore may be important in rumen and general bovine health.

**Fungi**

Rumen fungi ($10^3$–$10^6$ zoospores/ml) are anaerobic, falling into the class Neocallimastigomycetes, consisting of 6 previously recognised genera (*Anaeromyces, Caecomyces, Cyllamyces, Neocallimastix, Orpinomyces* and *Piromyces*) with 21 known species and, using molecular techniques, 2 recently discovered genera *Oontomyces*\(^{55}\) and *Buwchfawromyces*.\(^{56}\) Their discovery required a re-evaluation of the central dogma within microbiology that all fungi were aerobic.\(^{21}\) It has been proposed that these fungi evolved through
horizontal transfer of genes from bacteria. Anaerobic fungi may influence the rest of the microbial community as they produce H₂ during initial degradation of plant tissue and this H₂ can be used as fuel for the degradation mechanisms of other communities. Fungi colonize plant cell walls and account for approximately 8–12% of total microbial biomass in the rumen, but this figure varies widely depending on diet. Working with both the liquid and solid phase of rumen contents is important for the detection of fungal species as, while most plant fibre-associated fungi are retained in the solid phase, the liquid phase may contain smaller particulate matter that fungi may have attached to.

Although bacteria are the most prominent microorganisms in the rumen, fungi are the best degraders. They produce high levels of cellulases and hemicellulases, as well as possessing the ability to break down xylan due to the production of xylanases. Fungi appear to initiate the feed breakdown process, indicating that anaerobic fungi may be pivotal for feed utilization efficiency and animal growth and production in pasture-fed ruminants. As mentioned, anaerobic fungi are believed to have evolved through horizontal gene transfer from bacteria, and are the only known fungi to retain cellulosomes, which are cell-wall associated multienzyme complexes. Metagenomic approaches may be able to indicate the degree of horizontal gene transfer and where these genes were transferred from. By breaking down carbohydrates, fungi produce metabolites used for nutritional purposes by the host. They have many features which make them unique in comparison to fungi found outside the rumen, including polyflagellate zoospores (the reason why rumen fungi were originally mistaken for ciliate protozoa), hydrogenosomes, cellulosomes and a wide enzymatic spectra.

Bacteriophage

Bacteriophage are obligate pathogens of bacteria and occur in dense populations of approximately $10^7$ – $10^9$ particles per gram of digesta in the rumen. As is the case for other populations, bacteriophage abundances are also influenced by external sources, meaning they may also be controlled through different strategies. The bacteriophage and virus population found in a sample is referred to as the virome. The high number of rumen bacteriophage suggests that they may have an important function in the balance of the rumen system, but there is little known about the effect of the rumen virome on the system it inhabits. Viruses, however, have been shown to be a driving factor for evolution of many microbial systems in different environments, often facilitating horizontal gene transfer. Indeed Garcia-Vallve et al. showed that this was important in the transfer of glycoside hydrolase between bacteria and rumen fungi (Ricard et al.). proposed that this may also be important in the case of rumen ciliate protozoa, whereby horizontal gene transfer occurs from rumen bacteria in order to aid in protozoa adaption to the carbohydrate-rich environment of the rumen, transferring genes encoding plant cell wall degradation. This information indicates that bacteriophage may represent a shared gene pool for the rumen ecosystem. Again, metagenomics allows us to determine the origin of these genes. In a study carried out by, it was found that most viruses found in the rumen were associated with the most dominant phyla, namely Firmicutes, Bacteroidetes and Proteobacteria. Bacteria-phage interaction can be identified by the presence of clustered regularly interspaced short palindromic repeats (CRISPR) and CRISPR-associated proteins in the microbial population. These genes are fundamental in adaptive immunity in some bacteria and archaea, which enable the organism to respond and eliminate invading genetic material. Once infected by these invading genetic materials, the new DNA is integrated into the host CRISPR locus as new spacers, ultimately encoding a unique spacer sequence.

Techniques for analysis of the rumen microbiome

Culture-dependant approaches

The gastrointestinal tract of ruminant animals has a wide range of extremities, making it difficult to replicate conditions for optimal growth. While specific microbes can grow and can be characterised, a large percentage are unculturable in vitro and cannot be grown on laboratory media.
Culture-dependant techniques rely on various selective and enrichment culture conditions in order to replicate the microbes’ natural environment. Culturing anaerobes is quite difficult due to the need to exclude oxygen, the slow growth of the microbes and the complexity of other growth requirements (Rufener et al.).

65 used a continuous culture system in order to replicate the rumen environment. This technique, along with similar techniques, was used for the enumeration and identification of rumen microbes.

Using the solid phase of rumen contents can pose many problems when attempting to culture microbes. Many microbes adhere to particulate matter and are thus difficult to separate. Methylcellulose solution can be used to encourage detachment of bacteria from feed particles by providing a readily available feed source as described by Fessenden.

Traditional methods of classifying rumen bacteria were based on the standard bacterial identification methods; morphology, shape and Gram stain. Nutritional requirements and fermentation end products were also used as a means of classification. Roll tubes came to be employed to grow and isolate anaerobic species, and were used instead of conventional agar plates. The method was developed by Hungate,18 whereby agar medium was distributed in a thin layer over the internal surface of a test tube, which was then flushed with an oxygen-free gas of choice. According to Hungate, the tube required no particular incubation and could be examined with no anaerobic precautions necessary. The Hungate 1000 project, named after the aforementioned Robert Hungate, one of the first rumen microbiologists, seeks to combine culturing and high throughput sequencing approaches. More specifically, it is a catalogue of reference genomes from the rumen microbiome, which aims to produce a reference set of 1000 rumen genome sequences of cultivated rumen bacteria, ciliate protozoa, archaea and fungi. The project focuses on rumen bacteria and archaea, with a similar project, The Fungal 1000, focusing on fungal cultivation. The aim of the project is to support international efforts to develop methane mitigation strategies and rumen adaption technologies, as well as to initiate genome enabled research into understanding rumen function, feed conversion efficiency, methanogenesis and plant cell wall degradation in order to find a balance between food production and GHG. Along with this, the project will aid in discovering genes encoding fibre degradation that will enhance animal production and other functional genes such as those involved in bio-fuel production.20

Creevey et al.67 assessed several culture collections and databases, and found that members of the phylum Firmicutes and family Lachnospiraceae were the most commonly cultured bacteria from the rumen. Bacteriodetes represented just a small number with the strains identified belonging to just 2 genera. It was concluded that although the culture collections identified all of the major taxonomic groups that are widely known to be key players in rumen function, they do not represent the full diversity of the rumen microbiome. As stated previously Krause et al.,21 estimated that only 20% of the rumen microbiota can be cultured using standard techniques, and, more recently McCabe et al.,68 suggested that culturable microbes in the rumen account for less than 1% of the total microbial species.

**Culture-independent approaches**

Culture-independent methods, or, more specifically DNA-based methods of identification and detection of microorganisms, allow the examination of microbial communities at a molecular level. Metagenomic analysis allows the description of a microbial community by high throughput sequencing technology. Methods associated with this type of analysis include 16S rRNA and Internal Transcribed Spacer (ITS) amplicon sequencing, used for bacterial and fungal communities, respectively, or, shotgun sequencing, where DNA fragments are sequenced randomly, regardless of the microbe from which they came. Targeting of the mcrA gene has also been suggested in recent studies as a means of identifying methanogens.69 All of these methods use bioinformatics as a means of analysis and can be used in order to compare different microorganisms within a diverse ecosystem. Although these methods identify unculturable microbes, they do not provide a strain for further study. They may, however, provide insights that would allow the culturing of species that, thus far, have not been cultured. Understanding community structure is an important part in the
recognition of how microorganisms are affected by the environment which they live in and how they in turn affect the host organism. DNA sequencing technologies have transformed research of microbial and animal ecosystems. They have completely changed the approach involved in identifying microorganisms and the limitations, outlined above, associated with culture-dependant studies. Several platforms are available to achieve results; these differ only in small details, and all follow the same basic protocol of template preparation and clonal amplification, followed by rounds of parallel sequencing. Strategies used by each platform determine quality, quantity and bias of sequenced data. Figure 2, identifies the steps which may be included for metagenomic, metabolomic and proteomic studies.

16S rRNA amplicon sequencing requires the use of primers in order to identify the presence of specific bacteria and archaea. 18S target regions can also be used in identification of protozoa but may run the risk of amplifying bovine DNA, thereby affecting results. The ITS region found in fungi is the most widely used region to study the ecology of fungi. Various primers have been designed in order to yield the highest diversity. 16S sequencing is useful as it provides a good phylogenetic marker and there is a large database of sequences to assist in data analysis. It can identify highly conserved regions that are identical in all bacteria, with the use of a single primer pair. However, primer specificity is key and lack thereof will contribute to bias in detection of a target organism. Often, it is difficult to choose or design a primer that is specific to diverse groups, such as Firmicutes. Incorrect PCR
conditions, such as improper annealing temperatures during the amplification step in 16S, may result in false amplification. Additionally, 16S rRNA sequencing only gives information on bacterial populations, not fungal, viral, protozoa or amoeba, all of which are believed to play an important role in the rumen, as discussed earlier.

Kunin et al. \(^7\) defined metagenomics as the application of shotgun sequencing to DNA obtained directly from an environmental sample, producing at least 50Mbp of randomly sampled sequence data. Shotgun sequencing comprehensively samples all genes in all organisms, thus helping to explain what is there, what they are doing and to potentially examine why they are doing it. It evaluates total microbial diversity, detecting the abundance of microbes in a given environment. An important function is that it by-passes the need for obtaining pure cultures for sequencing, as this is often not possible for samples obtained from complex environments such as the rumen. It is a modern technology that is continuing to improve our ability to understand complex ecosystems, enabling the analysis of previously undetectable and unculturable microorganisms, otherwise difficult to analyse. It is also able to detect low abundance members of the microbial community, microbes which are dormant or inactive.

While the application of culture-based approaches to study ruminal contents has declined in recent years, our understanding of the rumen microbiome across different diets, species and geographical location has expanded due to the introduction of high throughput sequencing techniques. \(^6\) Culture-independent approaches rely on bioinformatics tools to handle large datasets. Programmes such as Kraken, assign taxonomic labels to short DNA sequences achieved through metagenomic studies. \(^2\) Superfocus, used in functional analysis of unannotated shotgun data (discussed below), classifies each sequence in the metagenome into a subsystem. \(^3\) Other programmes and databases used include: Qiime, RDP and Humann2. Statistical packages are then used to analyse the data.

**Dietary interaction with the rumen system**

**Feed conversion efficiency**

Feed conversion efficiency can be defined as kg of milk of standardised composition with respect to protein and fat concentrations produced per kg of dry matter consumed. \(^4\) It is an important measure of the performance of a farm feeding system. Residual feed intake (RFI) is a measure of feed efficiency, defined as the difference between the actual feed intake of an animal and its predicted feed intake based on maintenance energy requirement and growth rate. \(^5\) Many factors affect feed efficiency, with forages having the greatest effect due to making up the largest component of the slowly digestible portion of the diet. \(^7\) The amount of feed fed, and the ingredients contained in the ration, can influence feed efficiency. \(^8\) High quality, rather than poor quality, forage, also improves feed efficiency and is essential to farm profitability. Supplementary feeding in a grass-based system is necessary as nutritional needs cannot be met solely on a grass-based diet during times of the year when grass quantity and quality is not at its optimum. Improvements in feed efficiency would reduce on-farm costs and lead to production system efficiency. \(^6\) Supplementing forage with concentrates, such as cereal grains, improves feed efficiency, due to grains and their by-products breaking down more effectively in the digestive tract. As our understanding of the rumen microbiota deepens, and it’s associated effect on the host, it is possible to make links between microbial communities and host phenotypes. \(^9\) One of these being feed conversion efficiency. Metatranscriptomic analysis of the rumen microbial community has shown significant differences in both bacterial and archaean community structure between high residual feed intake groups and low residual feed intake groups. \(^8\) It was found that the microbiome of low RFI groups had the potential to adapt more
quickly to environmental stresses, such as that of intense dietary changes therefore. Changes in the abundances of the family *Lachnospiraceae* were noted between groups. Members of this family are associated with butyrate production, a VFA produced through the digestion of carbohydrates. The higher abundance of this family in high RFI groups may indicate an increase in butyrate metabolism, therefore, a resulting impact on feed efficiency. In contrast, the order *Methanomassiliicoccales* were more abundant in low RFI animals. Members of this order can use methyamine as an energy source, which may be harmful to the host, to produce additional ammonium during methanogenesis.

In high concentrate diets, methane reduction is induced via an increase in propionate which decreases H₂. Enteric methane emissions are also influenced by concentrate composition. Loss of energy in the ruminant digestive tract is a consequence of the production of methane. Therefore, including feeds which decrease methane production is important to reduce energy loss in the animal. Ruminal methane production represents approximately 2 – 12% of gross energy intake, which could otherwise be used for animal growth or milk production. Lowering emissions would benefit both the environment and the efficiency of livestock production. The interest in feed conversion efficiency is growing due to the need for more animal-derived protein and energy sources as well as minimizing the carbon footprint of the livestock sector.

Methane production through fermentation involves the conversion of almost all polymers to methane and carbon dioxide under anaerobic conditions. It is the result of a number of metabolic interactions among the microbial ecosystem described above. As reviewed by Sirohi et al., Wolfe began the first studies of CO₂ reduction to methane in the early 1970s, discovering six new coenzymes in the following years. He found that the process of methanogenesis requires seven coenzymes and eight enzymes. CO₂ and methane are removed by eructation to the atmosphere while acetate, propionate and butyrate travel across the rumen wall to the blood stream. Methane is produced by utilising simple substrates at low reduction potential to produce cellular energy. Methane synthesis contributes to the efficiency of the entire system and avoids the build-up of H₂, which would inhibit the normal function of microbial enzymes involved in electron transfer. Indeed, if ruminants did not produce any methane, the rumen pH would drop and fibre digestion would no longer be feasible.

A recent study suggested that methane production can, however, be reduced substantially without adverse effects on fibre digestion. The authors examined the effects of methane inhibition in the rumen of steers. The animals were fed either a hay:roughage diet or a hay:concentrate diet at a ratio of 60:40. An anti-methanogenic compound was also included, in this case, and in many others, chloroform. Results showed that with increasing levels of chloroform, there was an increase in H₂ expelled and CH₄ production decreased. Figure 3 illustrates the degradation of plant fibre from the hydrolysis of polysaccharides, releasing monomers. The monomers are then fermented through acidogenesis, resulting in the production of organic and short chain fatty acids. Hydrogen and CO₂ are also formed and are used by methanogenic archaea to produce methane. Animals on the hay only diet showed a more efficient redirection of H₂ into other microbial products in comparison to hay:concentrate diet. Metabolomic studies showed that there was an increase in the levels of amino acids, organic and nucleic acids found in the liquid phase of the rumen contents in both diets when methanogenesis was inhibited.

![Figure 3. Microbial methanogenic degradation of plant fibre in bovine dairy animals.](image-url)
This suggests that there may be enhanced microbial protein synthesis under these conditions. These changes showed an obvious alteration in the rumen microbiota, with an increase in the ratio of Bacteroidetes and Firmicutes and a decrease in Archaea and Synergistetes. However, there were no significant changes observed in the abundance of protozoa, fungi or fibrolytic bacteria, meaning that fibre degradation was not affected. The conclusion was made that, although there was a 30% decrease in methane production, this did not negatively affect rumen fermentation and fibre degradation, and the microbes were able to adapt and redirect H\textsubscript{2} to produce other end products. Further supplementation may be needed in order to drive excess H\textsubscript{2} and improve the energy supply to the animal. Although it had been believed that without the formation of methane, the rumen microbiota would be drastically affected, this study shows that it can adapt to changes in H\textsubscript{2} flux and produce different end products without the need to produce methane, in turn reducing methane eructed by the host animal.

There have been many attempts to minimise energy loss as a result of methane production, some of which have been mentioned earlier in this review. Many studies have explored the application of compounds and natural products to feed, prior to consumption by the animals. A novel, promising approach, is the use of probiotics. Probiotics present health benefits for the host when provided in sufficient amounts. explored the use of Bacillus licheniformis as a novel probiotic to reduce enteric methane emissions (Deng et al.). The study found that the use of the bacteria as a probiotic was sufficient in inhibiting methane emissions in sheep, increasing total tract digestibility and improve NUE. However, the subsequent effect on the overall rumen microbiome was not examined.

**Emerging areas of interest**

The usage of antibiotics in animal production systems has encouraged the abundance of antimicrobial resistance genes in farm environments to increase. It is a current threat to both human and animal health. Antibiotics are used therapeutically to treat or prevent specific diseases and those that are not entirely absorbed are excreted in the faeces and urine. As animal manure is collected in large pits for use as a high nutrient fertiliser, antimicrobial compounds can then be transferred to the soil. This may result in the possible uptake of these compounds by plants. Different land uses may require higher quantities of manure in order to provide higher concentrations of nutrients needed by specific plants. For example, land with which will be used for ensiling of grass may have larger quantities of manure applied in comparison to land used for seasonal grazing. This, in turn may result in varying abundances of AMR genes found in animals. In a study carried out by, the influence of diet on the ruminal resistome and abundance of pathogenicity genes was examined. The animals used in the study were antibiotic-free beef cattle on two different diets consisting of different levels of concentrate. Over 200 genes associated with antimicrobial resistance were detected across 4966 metagenomic genes, with results showing a higher diversity and abundance of these genes in animals fed a higher level of concentrate than animals on lower levels of concentrate. These results suggest, therefore, that not only does diet impact the microbial ecosystem but it also has the potential to impact antimicrobial resistance in the gut.

Interestingly, a recent study on the human gut microbiota, suggest that even non-antibiotic compounds, such as anti-inflammatory drugs somewhat overlap. This then poses the question as to whether non-antibiotic compounds could have a comparable impact on the gut microbiota of cattle.

**Emerging metagenomic techniques**

The rumen microbiota is a topic that has been covered in great detail since the emergence of NGS technologies. Most studies use reference databases such as Kraken (shotgun sequencing) and Greengenes (16S rRNA), among many others, to characterise the microbiota. The use of contigs or longer contiguous sequences for metagenomic analysis, may provide a deeper insight into the workings of the...
microbiome. Shorter reads, although having its advantages, do not represent complete genomes and may contain contamination. Studies have shown that the use of contigs may be an invaluable technique for the recovery of microbial genomes. Hess et al. (2011)\textsuperscript{98} were the first to use metagenomic binning techniques for the analysis of the rumen microbiota. Fifteen uncultured microbial genomes were assembled to > 60% completeness. The technique was also used in a study of the moose rumen microbiome, which recovered 99 metagenome-assembled genomes.\textsuperscript{99} The most recent study by Stewart et al. (2018),\textsuperscript{100} assembled 913 microbial genomes from metagenomic data of the rumen of cattle. This resulted in over 69,000 proteins thought to be involved in carbohydrate metabolism, in addition to the expansion of the Erysipelotrichales order. The study has, therefore, contributed immensely to improved metagenomic classification.

This study also highlights the potential to use hybrid short-read and long-read sequencing approaches in order to obtain complete end to end assembly of microbial chromosomes direct from metagenomic data.\textsuperscript{100} This approach may involve the use of both Illumina technologies for shorter reads and Pac-Bio or Oxford Nanopore MinIon for longer reads.\textsuperscript{101} The advantages of this approach include providing data on within-species genomic variation,\textsuperscript{102} identify pathogenic variants of species and determining the presence of pathogenicity islands and plasmids important for the detection of AMR.\textsuperscript{101}

**Protein metabolism**

Protein consumed by ruminants can be described as rumen degradable protein (non-protein and true protein N) or rumen un-degradable protein.\textsuperscript{50} Dietary protein intake determines the outcome of several factors associated with both the usable by-products and non-usable by-products, namely milk nitrogen efficiency, urinary nitrogen losses and ammonia emissions for the ruminant faeces.

**Nitrate/nitrite cycle**

The protein profile of feeds ingested by ruminants is extensively altered before reaching the small intestine due to fermentation within the rumen. This is primarily due to the conversion of nitrogenous compounds to microbial protein. The entire process is controlled by the fermentation of carbohydrates with recycling of nitrogen between the host and the gut.

The rumen has the ability to reduce nitrate to nitrite and nitrite to ammonia. Conversion of nitrate to ammonia leads to an increase of nitrogen available for use by the rumen microbes.\textsuperscript{103} It is thought that feeding nitrate to dairy cows may be an alternative method of reducing methane emissions by suppressing methanogenesis in the rumen. This is because, thermodynamically, nitrate and nitrite reduction to ammonia is a preferred reaction compared to the formation of methane with CO\textsubscript{2} as an electron acceptor, consuming electrons at the expense of methanogenesis. Recent studies have investigated the use of 3-Nitrooxypropanol (3NOP) to inhibit methane production.\textsuperscript{104,105} In a study carried out by,\textsuperscript{105} rumen methane emissions were decreased linearly by 30% through the use of 3NOP, while increasing milk protein and lactose yields. Animal performance also increased, suggesting that energy previously wasted through the production of methane was instead used by the host to increase body weight. It is also important to note that the inhibitory effect continued over the 12 week period of the trial, thus suggesting this is a means of effective methane mitigation which can be applied to the livestock industry. Although evidently an exceptional means of reducing methane emissions, the safety of 3NOP is questionable in terms of milk processing due to fears that excess nitrate passing from the rumen to the host could result in trace amounts of nitrate being found in the milk. Trace amounts are often not a concern but, with most milk being processed for use as milk powders, nitrate may become concentrated after drying stages, and even more cause for concern is the use of these powders for baby formulas where nitrate can cause methemoglobinemia or “Blue Baby Syndrome”. Therefore, a need for more comprehensive studies on milk produced by animals on 3NOP will need to be carried out in order to certify the safety of end products.
Nitrogen fractions in milk

The composition of milk is important in milk processing as it affects the quality of the product produced and the economic output of the dairy industry. Milk composition is an important factor with respect to milk processability. Key components are milk protein fractions, i.e., casein, whey and non-protein nitrogen (NPN), and minerals, with the most important being calcium, phosphorus and sodium. NPN refers to any compound that is not a true protein but can be converted into protein in the rumen following microbial synthesis. NPN is currently wasteful with regard to the processing of milk, thus a push towards having a larger proportion of casein or whey, relative to NPN, in the milk would mean more profitability for the processors per kg of milk produced. A high milk protein quality is key to maintaining thermal stability and gelation during processing. Urea is the most available NPN and is quickly broken down by bacterial ureases to form ammonia. The ammonia that is formed is important for bacterial growth, as it is used for amino acid synthesis, required for optimum growth. However, when ammonia is produced in high concentrations in the rumen, a certain amount is reabsorbed back into the bloodstream, converted back into urea in the liver and excreted through the kidneys and passed out as urine. It should be noted that urea may play an important role in low protein diets, compensating for the low concentrations of dietary protein in the diet, making urea utilization higher in comparison to that of high protein diets.

Nitrogen utilization efficiency

The importance of reducing nitrogen loss is critical to agriculture from a nutritional, environmental and economic perspective. Improving nitrogen utilization in dairy animals is fundamental to dairy product production and the environment. The agricultural industry contributes to approximately 40% of anthropogenic nitrous oxide emissions in Europe, with the dairy industry accounting for the most nitrogen loss in faeces and urine in comparison to any other ruminant animal in the agri-food sector. Not only does nitrogen loss have a huge impact on the environment, it also has huge consequences for the agricultural industry financially. Claimed that feeding nitrogen for optimum milk yields rather than maximum milk yields is the best way to reduce nitrogen pollution on dairy farms, without having consequences on productivity or profitability. As mentioned previously, dairy farms relying on grazed grass as their primary feed source may experience problems in improving efficiency of protein utilization due to the naturally high levels of protein in well-managed grass.

A means of improving nitrogen utilization efficiency (NUE) is by trying to synchronise the supply of energy and protein in the rumen through the use of dietary supplementation. Dietary protein and energy sources degrade in the rumen at different rates, available for use by the host at different times. Rumen synchrony refers to the provision of dietary protein and energy to the rumen so that they are available for use in proportions required or used by rumen microbes. Reid et al. examined the effect of dietary crude protein on grass-fed dairy cow production, nutrient status and milk heat stability. The study found that supplementing with high crude protein concentrates increased milk urea nitrogen concentration and milk non-protein nitrogen concentration, in comparison to supplementing with low crude protein concentrate. It was also found that increasing dietary crude protein lowered milk heat stability, which is undesirable in the processing of milk. Examining the nitrogen supply to the rumen based on the varying concentrations of protein consumed and how it effects the rumen environment and its microbial ecosystem is important in terms of determining what microbes can breakdown the feed stuff and use it efficiently, without wasting energy. Feeding animals more protein than is needed results in unnecessarily high feeding costs, reduced profits and further reductions in nitrogen utilization efficiency. High levels of dietary crude protein have a positive effect on protein degradation in the rumen, owing to increased ammonia concentrations, and have been shown to reduce nitrogen utilization efficiency for milk production. Reid et al. found that supplementation improved NUE in comparison to a grass only diet. Supplementation is important as it aims to provide further nutrients. Some studies have shown that supplementing with high crude protein concentrates may increase milk nitrogen efficiency, reducing nitrogen loss in faeces and urine, improving overall production efficiency, and reducing environmental impact.
for the animals that may not have been provided by grass alone.

**Conclusion**

In conclusion, the rumen microbial ecosystem is diverse and complex, consisting of microorganisms working symbiotically to break down feedstuffs consumed by ruminant animals. As addressed throughout the review, the microbiome controls the production efficiency of the animal, with certain pathways (such as those associated with methane production) resulting in energy loss in the animal. The microbiome also affects end-product quality (milk and meat) but also contributes to environmental pollution. High throughput sequencing-based techniques allow one to identify what microbes are there and, depending on the approach taken, what they are doing. Understanding the rumen microbiome and its connection to the ruminant itself is important for producing quality products, increasing profitability and reducing environmental impacts. Identifying certain metabolic pathways and further research into these pathways may determine the best diet for bovine ruminants in order to minimise energy loss, reduce methane production and increase nitrogen utilization efficiency. Examining the rumen microbiome can identify the effects of diet on the microbiome and in turn, the effects on milk yield, protein percentages, urea percentage (used as a NPN indicator) and milk protein yield. The composition of proteins found in milk can have a profound effect on milk processability, thus identifying pathways which may influence milk composition and improve processability is important.

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