PCR and qPCR-based applications in rumen microbiology research: a review

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ABSTRACT - The rumen and its microbial ecosystem play a central role in the overall nutrition and health of ruminant animals. However, development and homeostatic state of the entire gut system is influenced by different interrelated factors. Recent developments in molecular diagnostic tools by using amplicon sequencing of 16S ribosomal RNA and use of high-throughput data generated through applications of pyrosequencing is a promising approach to defining the rumen microbial genome. Several “omic”-based technologies based on PCR and real-time PCR (qPCR), are currently being used in gut microbiology in order to predict how the gut microbiota works. Such procedures include pyrosequencing, genome-wide shotgun sequencing, short metagenomics sequencing and metagenomics analysis, bacterial DNA integration for editing genomes of isolated microbes, etc. These tools are sensitive and precise in quantitation, identification and functional characterisation of the entire rumen microbiome. PCR/qPCR enables investigations of changes in the microbiome and microbiota with respect to age, diet, species and environmental variations thus providing new information about rumen microbial genome. In this review, we will highlight recent findings using PCR and qPCR-based procedures in investigating the complex nature of the rumen microbial population which has advanced our knowledge and understanding of the rumen microbial genome.

Keywords: Bacteria, Genome, Microbiome, Microbiota, Rumen, PCR/qPCR, 16S rRNA.

INTRODUCTION

One of the most widely used molecular diagnostic tool since its invention in the twentieth century is the polymerase chain reaction, PCR. PCR amplifies a segment of the DNA, such that millions or even billions of copies of a DNA molecule are generated in a very short time. PCR has been described as a versatile tool with diverse applicability to many different situations in molecular diagnostic research (Barlett and Stirling, 2003). It has also been used to detect DNA sequences, in forensic investigations, DNA fingerprinting, detection of bacteria and viruses to diagnose genetic diseases as well as cloning. PCR is the key that
unlocked several advances made in our understanding of the genome of hu-
man, bacteria and other species (Mullis et al., 1986; Innis and Galfand, 1990).

Bacteria, protozoa and anaerobic fungi have been implicated as key degraders of polysaccharides in the rumen (Schofield, 2000). They are found in all the four compartments of ruminants gut (rumen, reticulum, omasum and aboma-
sum). Microbiologically, the rumen is said to be made up of a complex ecosys-

tem which is composed of cells/ml concentrations of about $10^{10}$, $10^6$ and $10^4$
for bacteria, fungi and protozoa respectively, which interact in synergy and are active throughout the process of digestion and fermentation. This synergistic interaction also augment enzymatic digestion process in the animal for energy generation and yields volatile fatty acids (VFAs): acetate, butyrate, propionate, as well as other compounds such as ammonium, formic acid, methane gas, H$_2$, as well as CO$_2$ as end products of rumen fermentation (Krause et al., 2003; Pitta et al., 2010; Stewart and Bryant, 1988). Biomass-degrading enzymes encoded in the rumen microbiome and proteins belonging to bacterial and archaeal genomes have been associated with carbohydrate metabolism (Stewart et al., 2018).

Different bacterial groups function as one unit and assist the host (ruminants) degrade and utilize fibrous feed stuff in order to make volatile fatty ac-
ids as a source of energy from plant structural and nonstructural carbohy-
drates and proteins. Noteworthy, due to their relative abundance and meta-

bolic diversity, *Fibrobacter succinogenes*, *Ruminococcus albus* and *Ruminococ-
cus flavefaciens* (Hungate, 2013; Zhou et al., 2015) are main cellulytic bacteria associated with ruminal digestion of plant cell wall materials whereas *Rumi-
nobacter amylophilus* or *Prevotella ruminicola* dominates the degradation of starch (Steward and Bryant, 1988; Mobashar et al., 2019). Amino acids, soluble proteins and insoluble, particulate proteins are primarily degraded by bacteria and protozoa respectively (Cotta, 1998; Hino and Russel, 1987; Foroozandeh et al., 2009; Russell, 2009) while protozoa were reported to be limited in their ability to assimilate peptides (or amino acids) (Hino and Russel, 1987).

Classical culture-based procedures which are applied in isolation, identifi-
cation and quantitation of rumen microbes have been reported to account for less than 30% of the rumen microbial population. This is because several ru-
men microbes have shown tendencies of bypassing or escaping conventional culturing procedures and this necessitates the application of enhanced and more sensitive techniques (Nathani et al., 2013). However, recent develop-
ments in molecular diagnostic tools and use of high-throughput data generated
through applications of next-generation sequencing, pyrosequencing of bacterial 16S ribosomal RNA gene has provided wider understanding of the operations and activities of rumen resident microbes. Additionally, genome-wide shotgun sequencing for metagenomic data generation have revealed new information and innovative research areas about the function, characteristics and diversity of rumen microbiota. This has provided linkage between functional gene activity, metabolic pathways and rumen metabolites (Nathani et al., 2013; McCann et al., 2014; Roehe et al., 2016; Denman et al., 2018; Stewart et al., 2018). Research in rumen microbiome is quite complicated and consists of different areas which provides a wide range of research possibilities. Figure 1 shows some of these areas that are PCR/qPCR-based procedures in analysing, characterising or quantitating the microbial diversity in the rumen.

**Development of Rumen Microbiota**

Microbial genomes remain unchanged for life while the microbiome profiles tends to change as the new born animal grows (Goodrich et al., 2017). The rumen and its microbial ecosystem play a central role in the overall nutrition and health of the animal. However, development and homeostatic state of the entire gut system is influenced by different interrelated factors. These factors include genetic, physiological and environmental factors which may include dietary variations (quality and quantity), age, and maternal factors among others, are central in determining how the microbial population carryout their functions in supplying energy and nutrient to the animal as well as providing information about the variation in microbial diversity within the rumen microbiome (Callaway et al., 2010; Callaway et al., 2011; De Menezes., et al., 2011; Tymensen et al., 2012; Han et al., 2015; Dang et al., 2017).

Microbiologically, the rumen could generally be said to be made up of a complex ecosystem which is composed of cells/ml concentrations of about $10^{10}$, $10^6$, and $10^4$ for bacteria, fungi and protozoa respectively (Hungate, 1966; Skillman et al., 2006; Jewell et al., 2015), these microbes interact in synergy and are active throughout the process of digestion. Their synergistic interaction and effects tend to influence immunological responses, gut health and also augment enzymatic digestion process in the animal. As such, they are active in generating volatile fatty acids (VFAs): acetate, butyrate, propionate, and other compounds such as formic acid, methane gas, H$_2$, as well as CO$_2$ for utilisation by the animal in energy generation for cellular and metabolic processes. These groups function as one unit and assist the host (ruminants) in degrading and utilising fibrous feed stuff in a symbiotic relationship in order to make volatile
fatty acids as a source of energy from plant structural and nonstructural carbohydrates and proteins. The significance of the rumen microbiota in ruminant and human nutrition necessitates careful research in order to elucidate their roles and mode of action (Krause et al., 2003; Kamara, 2005; Pitta et al., 2010; Taschuk and Griebel, 2012; Rodríguez et al., 2015; Roehe et al., 2016).

Figure 1: Rumen microbiome project workflow chart (Modified from McCann et al., 2014)
Microbial Diversity in the Rumen

Rumen Bacteria

Rumen development has a significant effect on the microbial diversity of the entire gut system. Bacteria are the most dominant and diverse microbial community in the rumen. Flint et al. (2008) reported that bacteria are the most diversified groups of microbes in the rumen, approximately 95% of total microbiota. Subsequent studies involving rumen microbiology revealed a very large number of bacteria present in the rumen (up to about $10^{11}$ viable cells/ml) (Kim et al., 2011; Jami and Mizrahi, 2012b). Since new born ruminants generally have no functional rumen at birth, development and establishment of the ruminal microbiota still has various controversies by so many researchers (Hungate, 1966; Jewell et al., 2015). As the young ruminant grows, the species and bacterial community also experience a shift in composition (Li et al., 2012). *Fibrobacter succinogenes*, *Ruminococcus albus* and *Ruminococcus flavefaciens* were quantified using real-time PCR targeting 16S rDNA and the results showed that *R. flavefaciens* was slightly more dominant compared to *F. succinogenes* with the population of *R. flavefaciens* and *F. succinogenes* are significantly more than *R. albus* species (Mosoni et al., 2007).

Rumen Protozoa

Protozoa play a key role in the hydrogen transfer between microbial species and methane production within the rumen microbiome (Salonen et al., 2014). Ciliates have been reported to be the most abundant protozoa resident in the rumen of both domesticated and wild ruminants, contributing to fiber digestion, control of CHO fermentation process and the possibility of having a negative effect on protein metabolism. Until the application of PCR in rumen microbiological studies, rumen protozoa was quantified based on a microscopic count and this has shown that the most abundant genus accounting for about 95% of the total protozoal population in the rumen, is the genus *Entodinium*. However, comparing microscopic analysis with PCR based protocols has shown that there are several species and genera of protozoans which are resident in the rumen and are yet to be fully characterised (Regensbogenova et al., 2004; Skillman et al., 2006; Sylvester et al., 2009; Tymensen et al., 2012). This necessitates the need to intensively investigate protozoal populations in the rumen because they play significant role in fiber digestion and have negative effect on microbial protein turnover. However, they have been reported to have variation in the number of their rDNA copies per cell during a life cycle and also their cell sizes vary inter and intra-protozoal specie (Prescott, 1994;
Dehority, 2003; McSwansea et al., 2007). A major limitation of real-time PCR in microbial quantitation as well as protozoal analysis is that qPCR-based approach is more expensive than microscopic counts procedures. However, qPCR is more sensitive and more accurate with the ability to detect 1 to 10 million protozoal cells using genus specific primers that was developed by Skillman and co-workers for detection of Entodinium (Skillman et al., 2006).

**Fungi**

Fungi initiate mechanical and enzymatic break down of plant fiber to allow access for secondary metabolisers (Boots et al., 2013). Anaerobic rumen filamentous fungi form extensive interlocking rhizoidal systems. Accurate quantification and identification within this group of microbes by direct count using culture-based protocols are difficult, especially when considering that environmental samples contain both polycentric and monocentric species (Denman and McSweeney, 2006). Six genera, Neocallimastix, Piromyces, Anaeromyces, Caecomyces, Orpinomyces, and more recently Cyllamyces, have been recognized, while 18 species of anaerobic rumen fungi have been described on the basis of their thallus morphology and their zoospore ultrastructure (Ligenstoffer et al., 2010; Chaucheyras-durand and Ossa, 2014). However, investigation using PCR-based high-throughput sequencing technology has revealed several uncultured taxa and the relative abundance of the rumen fungal populations estimated to be 10% of the total microbial biomass, with large variations according to diet and individual (Krause et al., 2013).

**Rumen Viromes and Bacteriophages**

The occurrence of bacteriophages in the rumen has previously been documented at $10^7$ to $10^9$ particles per ml. About 28,000 different viral genotypes were identified using PCR-based protocols. Despite the possibility of isolating and storing several rumen phages in culture collections, particularly in the 1970's and 1980's, it was reported that only those phages with applicability in genetic engineering and phage therapy were successfully characterised beyond an initial assessment on the basis of morphology (Gilbert and Klieve, 2015). DNA sequencing and the advent of metagenomic studies to comprehensively sequence phage particle fractions obtained from rumen fluid as well as the full extent of viral diversity within the rumen based on “omic” applications is gradually being revealed (Berg Miller et al., 2012). Thus, diverse groups of rumen resident phages were found to have a high tendency of infecting various
species of rumen bacteria through their tendency for exhibiting lytic replication (Gilbert et al., 2017). However, little information is available about the genetics of phages while their genome has remained largely uncharacterised. Gilbert et al. (2017) reported that bacterial genome sequencing project is a powerful tool which is revealing the presence and activity of phages and their ability to integrate their DNA into the genome of their host to form stable, lysogenic associations. Thus, rumen phages could infect and replicate within the host and the release of progeny phage particles (Gilbert et al., 2017).

Exploring the rumen microbiome using PCR/qPCR

**End point PCR**

Polymerase chain reaction, PCR, is a step-by-step in vitro primer interposing DNA enzymatic amplification process. It is a technique of making millions of copies of a particular DNA target of interest that is being replicated during PCR cycles over a period of time (Ginzinger, 2002; Sluijter et al., 2006). At the last cycle of the reaction, PCR products are detected and analysed using electrophoresis. The amplification process (detection and identification of target DNA) occurs in three stages; denaturation, annealing and elongation stages. Identification is made through visual methods based on size the amplified DNA piece (Freeman et al., 1999; Ginzinger, 2002; Sluijter et al., 2006). Microbial PCR procedures have been optimised using primers targeting V2–V4 regions of the 16S rRNA gene. Recently, pathophysiological examination in sheep by comparing resident microbiome in the upper aerodigestive and lower respiratory tracts of lambs, PCR was used to amplify the V2-V3 region of the 16S rRNA gene and subsequently sequenced via Illumina Miseq which revealed oropharyngeal swabs were either dominated by bacteria commonly associated with the rumen or by bacteria commonly associated with the upper aerodigestive tract (Glendinning et al., 2017). Similarly, PCR procedures have been optimised using primers targeting the V4 and V3-V4 region of the 16S rRNA gene under 20 and 28 PCR identical cycle conditions for amplicon sequencing to generate a relatively accurate representation of rumen microbiome (McGovern et al., 2018).

**RT-PCR; qPCR**

Real time PCR is also known as quantitative PCR (qPCR), depending on the application. The greatest advantage real-time PCR reaction compared to end-point PCR is the possibility to determine concentration of the targeted DNA fragment as the template DNA is being synthesized. This makes data to be collected throughout the PCR process providing the possibility to view the entire
reaction and product generation, also combining the amplification and detection in one step (Suzuki et al., 2000; Rasmussen, 2001; Wong and Medrano, 2005; Sluijter et al., 2006; Yibing, 2012; Alvarez and Doné, 2014). In qPCR, a major feature is the point during amplification at which PCR product accumulation is first detected instead of quantitation of PCR product after a defined number of cycles. Results from real-time PCR can be both quantitative and qualitative. In the latter case, it is possible to visualise the melting curve of a DNA of interest after amplification, which can prove the presence or absence of the gene of interest and specificity of the reaction and not the amount of DNA present in a given sample (Sluijter et al., 2006; VanGuilder et al., 2008; Malmuthuge et al., 2014; Glendinning et al., 2017).

Rumen microbial composition consist of bacteria, archaea, ciliate protozoa, fungi, bacteriophage and viruses. Over 200 species of bacteria (approximately $10^{10} - 10^{11}$ cells/ml) and 100 species of protozoa and fungi are found in the rumen. However, despite their abundance and significant roles in fermentation and microbial digestion in the rumen, it was difficult to have them cultured, identified or characterised using conventional culture-based methods. There are many several species and groups of bacteria, protozoa and fungi which tend to influence different physiological processes such as *Fibrobacter succinogens*, *Lactobacillus ruminis*, *Escherichia coli*, anaerobic chytridiomycete fungi, *Peste des petits ruminants*, rinderpest, etc in ruminants (Chaucheyras-durand and Ossa, 2014; Bainbridge et al., 2016; Mathew et al., 2018). These identified microbial species could vary with diet, genetics and environment (Benson et al., 2010; Uyeno, 2010; Kim et al., 2012; Henderson et al., 2015). The use of PCR and qPCR in microbial investigations have evinced that the abundance of rumen bacteria is up to about 1012 cell/ml (Kim et al., 2011; Jami and Mizrahi, 2012b). With real-time PCR, the number of isolated bacterial species has been significantly increased and was found to play salient roles in the rumen (Denman et al., 2018). Similarly, 16S rRNA-gene-targeted specific primers for analysis of caecal microbial community using qPCR revealed how dietary supplements and age influenced amount of the copy number of each bacterium (total bacteria, *Bacteroides* and *Clostridia*) found in the caecum (Bagóné Vántus et al., 2018). A peculiar feature of the real-time PCR is that the bacterial and total microbial populations can be measured concurrently, this is important when dealing with heterogeneous rumen samples (Skillman et al., 2006).

Application of optimised and sensitive protocols based on the use of PCR/qPCR has identified and characterised several rumen microbes which
have shown tendencies of by-passing or escaping conventional culturing procedures thus providing accurate and more detailed information about the nature of microbial diversity and interaction that exist within and between rumen microbial populations (Nathani et al., 2013).

**Bacterial Diversity:**

Phylogenetic analysis of the bacterial communities that colonise the gut system of pre-weaned calves by Malmthuge et al., (2014), revealed that a total of 83 genera belonging to 13 phyla with *Firmicutes*, *Bacteroidetes*, and *Proteobacteria* predominating the gastro intestinal tract of the calves under investigation. Subsequently, qPCR analysis of the selected dominant bacterial genera (*Prevotella*, *Bacteroides*, *Lactobacillus*, and *Faecalibacterium*) in the same study revealed that, the occurrence of the identified bacterial genera varies significantly with different locations of the gastro intestinal tract as well as between bacteria found on the mucosa and in the rumen digesta (Malmthuge et al., 2014). These findings were consistent with Henderson et al. (2015), who used a qPCR-based sequencing procedures and reported that *Bacteroidetes*, *Firmicutes*, and *Proteobacteria* constitute the most dominant bacterial phyla in the rumen while subsequent phylogenetic analysis of the genomic sequences showed that *Prevotella*, *Butyrivibrio*, and *Ruminococcus*, as well as unclassified *Lachnospiraceae*, *Ruminococcaceae*, *Bacteroidales*, and *Clostridiales* are a “core bacterial microbiome” at the genus level (Henderson et al., 2015). Similarly, data from pyrosequencing of the bovine rumen bacterial 16S ribosomal rDNA gene sequences evince that the average composition of the rumen bacterial community consisted mainly of the phyla *Bacteroidetes* and *Firmicutes*, 51% and 43% of all reads, respectively while *Proteobacteria* accounted for 5.455% of the reads (Kim et al., 2011; Jami and Mizrahi, 2012). Molecular diversity of methanogens in the rumen of goats investigated with 16S rRNA gene clone libraries which consist of rumen fluid-associated methanogens (FAM) and rumen particle-associated methanogens (PAM) using methanogen-specific primers. The data obtained from the qPCR results shows that there is a significant difference in the concentration of methanogens in the two investigated groups, PAM (8.97 log 10) > FAM (7.57 log 10) (Gu et al., 2011). Recently, several bacteria has been identified which are significantly more abundant in lung fluids than in the upper aerodigestive tract swabs in sheep; *Staphylococcus equorum* was reported to be most predominant (Glendinning et al., 2017).
Protozoal Diversity:

Real-time PCR is more sensitive than microscopic count when it comes to protozoal detection and quantification because with qPCR, up to about 1-10^6 cells/ml protozoa is detectable (McSwasea et al., 2007). Cell counts and real-time PCR has been applied in quantifying total rumen protozoa and different genera of rumen protozoa on the basis of morphology (microscopy) or primer choice (real-time PCR) (Skillman et al., 2006). Diversity of the rumen protozoa using protozoa-specific PCR primers revealed that major protozoal species found in the rumen includes Epidinium caudatum, Entodinium caudatum, and Isotricha prostoma (Sylvester et al., 2004). Tymensen et al. (2012) compared protozoan communities from hay-fed cattle and silage-grain-fed cattle using T-RFLP analysis and the outcome yielded similar overall results to data obtained from microscopic analysis; it was concluded that Entodinium was the predominant genus of rumen protozoa present in all cattle regardless of diet, accounting for 94.2% of protozoa in silage/grain-fed cattle versus 57.4% for hay-fed cattle (Skillman et al., 2006; Tymensen et al., 2012). Apart from Entodinium, which has been reported to be dominant in the rumen, large phylogenetic diversity was found from a few numbers of sequenced clones. Protozoa that inhabit the rumen were detected by PCR using protozoan-specific primers based on the 18S rDNA sequences in the rumen contents of cow and the predominant protozoan genus identified in the whole rumen belonged to the Entodinium group (81.1%) (Karnati et al., 2003; Shin et al., 2004). Entodinium accounted for about 98% of the total protozoa, and populations within the same sheep were relatively stable, but greater variation occurred among individual sheep (10^0 and 10^6 Entodinia per gram of rumen contents).

Fungal Diversity:

Earlier attempts using the PCR/qPCR in order to quantitate rumen fungi has shown that there is high level of conservation within fungal 18S rDNA gene sequences and a more appropriate target for identification known as the internal transcribed spacer 1 region (ITS1) has been proposed to be targeted. ITS1 is located between the 18S rDNA and 5.8S rDNA genes and this region. It was identified as containing high levels of sequence variation, and it is used for the phylogenetic identification of anaerobic rumen fungi using qPCR-based techniques (Bowman et al., 1992; Li and Heath, 1992; Brookman et al., 2000; Edwards et al., 2008). Phylogenetic diversity of the gut anaerobic fungi was investigated in 30 different herbivore species by Liggenstoffer et al. (2010) using the
internal transcribed spacer region 1 rRNA in addition to a total of 267,287 sequences representing all known anaerobic fungal genera were obtained. The study revealed that sequences related to the genus Piromyces are predominant in rumen, which is about 36% of the total sequences obtained. Other sequences were related to the genera *Cyllamyces* and *Orpinomyces* and were categorised as the least abundant, representing 0.7 and 1.1% of the total sequences obtained, respectively. Subsequently, 38.3% of the sequences obtained did not cluster with previously identified genera and formed 8 phylogenetically distinct novel anaerobic fungal lineages (*Liggenstoffer et al., 2010*). qPCR also revealed significant reduction in the relative abundance of fungi with respect to diet and dietary additives (*Tapio et al., 2017*).

**Rumen Bacteriophage:**

Bacteriophages have significant role in maintaining rumen microbial balance (*Berg Miller et al., 2012*) and assisting the progress of horizontal gene transfer in the rumen microbiome (*Rohwer et al., 2009; Berg Miller et al., 2012*). There is an association between bacteriophage and microbial populations in the rumen evinced by sequence similarities that between the rumen viromes and the rumen microbiome. The application “omic”-based procedures have shown that most abundant rumen bacteriophage and prophage have been reported to be closely associate with several members of the rumen dominant bacterial phyla, *Firmicutes*, *Bacteroidetes*, and *Proteobacteria* (*Brulc et al., 2009; Berg Miller et al., 2012*). *Berg Miller et al., (2012)* reported that total DNA isolation from a phage-enriched fraction of rumen fluid and subsequent pyrosequencing of the total rumen virus-enriched metagenome (virome) revealed that rumen microbial genomes tend to share some sequence similarity to rumen viruses (phages) and prophage which also agrees with earlier findings which suggests transfer of genetic information between the microbial and viral populations within the rumen (*Brulc et al., 2009*) while prophages have been reported to be more numerous than lytic phages by approximately 2:1(*Berg Miller et al., 2012*).

**Taxa Specific Oligonucleotide (Primer) Sequences for PCR**

Application of real-time PCR in rumen microbiology using species-specific bacterial PCR primers (*Table1*) have been developed. Each pair of PCR primer sequence targets the 16S rRNA gene for detection and quantification of bacteria from diverse species within the rumen microbiome (*Khayalethu, 2013*). Also, these primer sets have been applied in studying the bacterial variation of microbial population based on DNA copy numbers in relation to changes in diet,
Application of “omic” procedures with PCR/qPCR to rumen microbial diversity

Application of PCR/qPCR “omic”-based procedures has enhanced investigation of rumen microbiome over the past few years. Whole-genome sequencing, pyrosequencing, proteomics (metagenomic-proteomic), transcriptomics has provided a clearer insight into composition, functionality and diversity of each rumen microbial species (Morrison et al., 2003; Roesch et al., 2007; Dowd et al., 2008; Brulc et al., 2009; Callaway et al., 2010; Iakiviak et al., 2011; Ransom-Jones et al., 2012; Krause et al., 2013). Associations between microbial genes and the host animal could provide information on microbiome genotype-environment interaction using genome wide association studies, GWAS, in order to predict genetic predisposition to disease or performance among species (Goodrich et al., 2017). Such DNA technologies are the basis of current rumen microbiological research and are revealing information about the true nature of the rumen microbiome. Basic concept underlying analysis of rumen microbiome on the basis of taxonomic structure, diversity, function, and sequence analysis methodology has been described by the rumen microbiome project workflow shown on Figure 1 (Di Bella et al., 2013; McCann et al., 2014).

Metagenomics and Metatranscriptomics

Estimation of functional and fractional potentials of DNA and RNA can be achieved through the application of metagenomic and metatranscriptomic analysis. Such analysis are carried out using bioinformatics tools in order to study genetic materials from uncultured microorganisms. These tools have proven to be efficient in characterising the rumen microbiota according to function on the basis of their genomes as well as high throughput analysis of amplified taxonomic marker genes (Li et al., 2017). In metagenomics, the template DNA is sequenced without prior amplification of specific genes which results in a snapshot of the gene pool and functional potential of the microbiome while in transcriptomics, mRNA is analyzed to provide a measure of gene expression within the intestinal microbiome (Suchodolski, 2012). A deeper and clearer insight into taxonomic and functional characteristics of the rumen mi-
crobiome and its interaction with the environment offers researchers the opportunity to optimize the digestion process in the rumen for enhanced and efficient sustainable utilization of dietary nutrients (Walsh et al., 2017).

Metagenomic and metatranscriptomic inventions are considered more accurate and are widely used with conventional PCR and qPCR based protocols for routine identification, for result justifications and taxonomic assessment of the microbiota (Urich et al., 2008; Hong et al., 2009; Huber et al., 2009; Ross et al., 2012). The use of 16S rDNA sequences generated in metagenomics datasets sequencing procedures based on specific targeted marker genes and transcripts using 16S bacterial and archael rDNA/rRNA, 18S rDNA/rRNA for protozoa, and internal transcribed spacer (ITS) gene/transcript for fungi has been reported to be the basis of classification and phylogenetic analysis of rumen microbiome (Deng et al., 2007; Ellison et al., 2014).

Metagenomics and metatranscriptomics are fast, reliable, less expensive with diverse applicability. It has been applied in soils, water and animal-based samples. However, as a result of the low fraction of 16S rDNA reads present in metagenomics datasets, most metagenomic studies rely on analogous DNA amplicon sequence to characterise microbial populations (Urich et al., 2008; Baker et al., 2013; Martínez et al., 2013 Franzosa et al., 2014; Logares et al., 2014; Mason et al., 2014; Rooks et al., 2014; Tveit et al., 2014; Li et al., 2016). To characterise biomass-degrading genes and genomes, 268 gigabases of metagenomic DNA from rumen microbes (which are viscid to plant fiber incubated in cow rumen), Hess et al., identified 27,755 putative carbohydrate-active genes and 90 expressed candidate proteins, of which 57% were enzymatically active against cellulosic substrates (Hess et al., 2011). However, recent studies reported that the rumen microbes are under-represented in the public databases (Li et al 2017; Stewart et al., 2017; Stewart et al., 2018). From these recent studies, 220 high quality bacterial and archael genomes assembled directly from 768 gigabases of rumen metagenomic data were presented and the comparative analysis of these sequences against current publicly available genomes shows that over 90% of these sequences represent previously non-sequenced strains and species of bacteria and archaea. The genomes contain over 13,000 proteins predicted to be involved in carbohydrate metabolism in the cow rumen. Furthermore, rumen virome has been explored using metagenomics and metatranscriptomics in dairy cattle and the results have shown that the rumen virome is composed of highly diversified and vast number of phages (Berg Miller et al., 2012).
Table 1
PCR Primer sequences for detection of rumen bacteria, arachea and methanobacteriales

<table>
<thead>
<tr>
<th>Target bacterium</th>
<th>Primer sequence (5′–3′)</th>
<th>AT (°C)*</th>
<th>Ps (bp)**</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total bacteria</td>
<td>Forward: CCTACGGAGGCGACGAG</td>
<td>60</td>
<td>194</td>
<td>(Muyzer et al., 1993; Misoni et al., 2007)</td>
</tr>
<tr>
<td></td>
<td>Reverse: ATTACCCGGGCTGCTGG</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>*Streptococcus bovis</td>
<td>Reverse: CTAATACGGCATAACAGCAT</td>
<td>57</td>
<td>869</td>
<td>(Tajima et al., 2001)</td>
</tr>
<tr>
<td></td>
<td>Forward: AGAAACTCCTATCTCTAGG</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>*Eubacterium ruminantium</td>
<td>Forward: GCTTCTGAGAAATCATTGAGAG</td>
<td>57</td>
<td>671</td>
<td>(Tajima et al., 2001)</td>
</tr>
<tr>
<td></td>
<td>Reverse: TGTGCTTCAGTGCTGAGT</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>*Fibrobacter succinogenes</td>
<td>Forward: GTATGGAGTAGCTTGAAG</td>
<td>60</td>
<td>446</td>
<td>(Tajima et al., 2001; Wanapat &amp; Cherdthong, 2009)</td>
</tr>
<tr>
<td></td>
<td>Reverse: GCTGCCCCTGAATCAGTC</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>*Ruminococcus albus</td>
<td>Forward: CCCTAAAGCAGCCCTAGTTGAG</td>
<td>60</td>
<td>175</td>
<td>(Koike &amp; Kobayashi, 2001; Khampa et al., 2006)</td>
</tr>
<tr>
<td></td>
<td>Reverse: CTTCTTGCAGGCTAGTTGAG</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>*Ruminococcus flavefaciens</td>
<td>Forward: GGACGATAATTGACGGTACTT</td>
<td>62</td>
<td>835</td>
<td>(Tajima et al., 2001)</td>
</tr>
<tr>
<td></td>
<td>Reverse: GCAATCGAAGCCTGGCACAT</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>*Ruminobacter amylophilus</td>
<td>Forward: CAACAGTCGATTCAGA</td>
<td>57</td>
<td>642</td>
<td>(Tajima et al., 2001)</td>
</tr>
<tr>
<td></td>
<td>Reverse: CACTACTCATGGCACAT</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>*Prevotella bryantii</td>
<td>Forward: AGTCGGAGCGTGAAGATTG</td>
<td>68</td>
<td>540</td>
<td>(Tajima et al., 2001; Wanapat &amp; Cherdthong, 2009)</td>
</tr>
<tr>
<td></td>
<td>Reverse: CAAAGCGTTCTCAGACTC</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>*Prevotella ruminicola</td>
<td>Forward: GGTATCTCTGAGTGAGAT</td>
<td>53</td>
<td>485</td>
<td>(Tajima et al., 2001; Wanapat &amp; Cherdthong, 2009)</td>
</tr>
<tr>
<td></td>
<td>Reverse: CTGATGCAACATTGAAA</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>*Anaerovibrio lipolytica</td>
<td>Forward: TGGGTGTAGAATGGAGTTAC</td>
<td>57</td>
<td>597</td>
<td>(Tajima et al., 2001; Wanapat &amp; Cherdthong, 2009)</td>
</tr>
<tr>
<td></td>
<td>Reverse: CTCTTGCAGTAAGAAATAC</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>*Ruminobacter amylophilus</td>
<td>Forward: CAACAGTCGATTCAGA</td>
<td>57</td>
<td>642</td>
<td>(Tajima et al., 2001; Wanapat &amp; Cherdthong, 2009)</td>
</tr>
<tr>
<td></td>
<td>Reverse: CACTACTCATGGCACAT</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>*Selenomonas ruminantium</td>
<td>Forward: TGCTATTACCGGAATGTTG</td>
<td>57</td>
<td>513</td>
<td>(Tajima et al., 2001; Wanapat &amp; Cherdthong, 2009)</td>
</tr>
<tr>
<td></td>
<td>Reverse: TCCTGACTCAAGAAAAGA</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>*Treponema bryantii</td>
<td>Forward: ACTGCAGCGGACTGTCAGA</td>
<td>57</td>
<td>412</td>
<td>(Tajima et al., 2001; Wanapat &amp; Cherdthong, 2009)</td>
</tr>
<tr>
<td></td>
<td>Reverse: ACCTTACGGGTCAGTCTC</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total arachea</td>
<td>Forward: GYGCAGCAGGCCAGCTGTTG</td>
<td>145–420</td>
<td></td>
<td>(Takai and Horikoshi 2000)</td>
</tr>
<tr>
<td></td>
<td>Reverse: GGACTACCCGTTACCTTAAT</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Methanobacteriales</td>
<td>Forward: GGAAGGAGCTTGAAGCTGTC</td>
<td>343</td>
<td></td>
<td>(Yu et al., 2005)</td>
</tr>
<tr>
<td></td>
<td>Reverse: TACGGTGCCACTGCTT</td>
<td></td>
<td></td>
<td></td>
</tr>
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</table>

*Annealing Temp., **Product size
Table 2

<table>
<thead>
<tr>
<th>Target microbe</th>
<th>Primer sequence (5’–3’)</th>
<th>Ps (bp)*</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td>General anaerobic fungi</td>
<td>Forward: GAGGAAGTAAAGTGCAACAAAAGGTATTCT</td>
<td>120</td>
<td>(Denman et al., 2006)</td>
</tr>
<tr>
<td></td>
<td>Reverse: CAAATTCACAAAGGGTACAGTATT</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Neocallimastigales-specific primer</td>
<td>Forward: TTGACAAGTGACCTCTTGGTTCCT</td>
<td>-</td>
<td>(Edwards et al., 2008)</td>
</tr>
<tr>
<td></td>
<td>Reverse: GTGCAATATGCGTGTTCTGCAGT</td>
<td></td>
<td></td>
</tr>
<tr>
<td>General Protozoa</td>
<td>Forward: CAYGTCTAGTGATAAAATAACTAC</td>
<td>-</td>
<td>(Sylvester et al., 2004)</td>
</tr>
<tr>
<td></td>
<td>Reverse: CTCTAGTGATTTTTAC</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ciliate Protozoa</td>
<td>Forward: TGTCCTGGTTAATTCCGA</td>
<td>-</td>
<td>(Isaq et al., 2014)</td>
</tr>
<tr>
<td></td>
<td>Reverse: GTGATRGTTTACTT</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*Product size

Additionally, cow rumen metatranscriptomic datasets revealed distinct taxonomic and functional signatures in a study by (Jiang et al., 2016) with enzymes involved in amino acid, energy and nucleotide metabolism and also identified microbiome-specific pathways such as phosphonate metabolism and glycan degradation pathways in the rumen. Rumen metagenomic profiling has been used to investigate relative abundance of microbial genes in the gastrointestinal tract of ruminants and could be applied to different species in order to predict influence of microbial composition on traits. Roehe and co-workers (2016) investigated the link between microbial genes, methane emissions and feed conversion efficiency in ruminants using metagenomic analysis and concluded that host microbial composition is an additional reliable basis for selection in animals since the host animal controls its own microbiota (Roehe et al., 2016).

**Bacterial 16S rRNA and Next Generation Sequencing (NGS) applications**

Bacterial 16S rRNA gene is common and conserved among bacteria species but vary between taxa and as such, PCR/qPCR-based protocols target the 16S rRNA for identification, classification or characterisation of bacteria since decades of PCR application in molecular microbiological procedures (Gutell et al., 1985; Noller, 1984; Ginzinger et al., 2002; Khayalethu, 2013). Next-generation sequencing (NGS) provides high-throughput data for precise and accurate analysis, characterisation and an in-depth examination of the complex rumen microbiome (Shokralla et al., 2012). NGS-based 16S rRNA sequencing is considered cost-effective technique for identifying microbial strains that may not be easily characterised using culture-based methods. Early studies by Gray et al., (1984) highlighted that the bacterial 16S gene is composed of about nine hypervariable regions (V1-V9) that ranged from about 30-100 base pairs long...
that are involved in the secondary structure of the small ribosomal subunits. However, the degree of conservation varies widely between hypervariable regions, with more conserved regions correlating to higher-level taxonomy and less conserved regions to lower levels, such as genus and species. Consequently, amplicon sequencing revealed diet specific taxa abundance variations revealing significant differences in protozoal and fungal composition within the rumen of a mature cow (Tapio et al., 2017).

Several studies have explored the sequencing of the 16S rRNA gene in evaluating the rumen epithelial bacterial diversity during the development of the rumen and the most dominant phyla detected in the rumen epithelium were found to belong to Proteobacteria, Firmicutes, and Bacteroidetes (Steveson & Waimer, 2007; Li et al., 2009; Petri et al., 2013; Chaucheyras-durand and Ossa, 2014; jiao et al., 2015). Additionally, Bacteroidetes has also been reported to be the most abundant phylum in the rumen microbiota, representing up to 74.8% of the 16S sequences, followed by Firmicutes (12.0%), Proteobacteria (10.4%), Verrucomicrobia (1.2%) and Synergistetes (1.1%) and the researchers concluded that the four most abundant phyla in the rumen of matured bull calves which were fed a solid diet are Bacteroidetes (52.0%), Firmicutes (42.7%), Spirochaetes (2.3%) and Fibrobacteres (1.9%) (Li et al., 2012).

**Pyrosequencing, Illumina Sequencing (MiSeq) and Massively Parallel Signature Sequencing (MPSS)**

Pyrosequencing procedure also enables comparison between and within bacterial species in a given rumen microbiome using 16S rRNA hypervariable regions (approx. 1500 base pairs long) as standard and reliable markers for the taxonomic classification and phylogenetic analysis (Yang et al., 2016). Pyrosequencing is a well-optimized procedure for phylogenetic analysis of the complex rumen microbiome (Kim et al., 2011; Klindworth et al., 2013; Yang et al., 2016; Fuks et al., 2018; McGovern et al., 2018). Tewari et al. (2011), showed that utilisation of sequence data from 2 variable regions within the 16S rRNA gene: V1 and V6 identified about 80% of microbes isolated from animals up to the genus level. Similarly, pyrosequencing of hypervariable V3-V5 regions of the 16S rRNA gene and whole-genome shotgun approach to rumen microbiota in pre-ruminant (42-day-old) calves has identified fifteen bacterial phyla in a study conducted by Li et al. (2012). Previously, it has been shown by Tewari et al. (2011) that the differences among different bacterial strains were identified using pyrosequencing. From the strain differential studies, they pointed out that Staphylococcus aureus differs from other Staphylococci while Rhodococcus...
equi also differs from the closely related genus Dietzia spp. Subsequently, pyrosequencing of S. bovis identified and differentiated the species from Enterococcus cecorum (Tewari et al., 2011). These findings are consistent with other studies which evinced that deep amplicon sequencing of the 16S ribosomal RNA gene have significant differences within the bacterial communities on dietary basis and between rumen solid and liquid contents (Callaway et al., 2010; Callaway et al., 2011; De Menezes, et al., 2011).

Similarly, taxonomic analysis of metagenomic reads from 16S rRNA sequences during investigation of the establishment of gut microbial community indicated that the predominant phyla were distinct at different growth stages (Han et al., 2015). They show that phyla Firmicutes and Synergistetes were predominant in rumen samples taken from 80 to 100-day-old goats. The age of the animals has shown to have significant effect on the abundance of bacterial species. Goats on day 100 showed Bacteroidetes and Firmicutes as most abundant phyla while relative abundance of Firmicutes and Synergistetes was found to be reduced after weaning and Bacteroidetes and Proteobacteria increased with increase with age (from 80 to 110 days). Similarly, Illumina MiSeq platform was used for amplification of the V4 hypervariable region of the 16S rRNA gene from rumen fluid of goats fed the same diet for 20 days. Analysis of the gene sequences using the quantitative insights into microbial ecology pipeline software shows that Bacteroidetes, Firmicutes, Tenericutes were the predominant phylum in both groups, and their relative abundance was 60.63%, 29.48%, 2.24% respectively. Prevotella being the most abundant shared genus between the control and experimental groups (Wang et al., 2018).

Massive parallel signature sequencing (MPSS) procedure is a tool that is used for an in-depth profiling of gene expression pattern through sequencing mRNA transcripts. Bioinformatics tools sort out sequence signature of about 16-20 base pairs generated and identified from each bead by counting the number of individual mRNA molecules produced from each gene (Reinartz et al., 2002). Thus the level of expression of each gene is directly proportional to the number signatures for the gene’s mRNA counted. It is highly specific, sensitive (with a level of sensitivity as low as a few molecules of mRNA per sample), and transcripts are captured on individual microbeads through a complementary DNA signature sequence without prior identification and characterization of the gene. This provides wide-range analysis of DNA templates transcribed into an RNA of an entire genome (Brenner et al., 2000; Reinartz et al., 2002; Jongeneel et al., 2005; Rédei, 2008).
Application of MPSS has also provided deeper insight into rumen microbiological research in recent years. It has possibility of providing accurate predictions that are based on the host genome for traits which are largely influenced by the gut microbiota such as digestive tract disorders, metabolic functions, body mass index (BMI), inflammatory bowel disorder (IBD) and enteric methane production in cattle (Leahy et al., 2013; Ross et al., 2013a). It could also be applied in genomic predictions of complex phenotypes in humans and animals (Ross et al., 2013a). Diversity of the rumen virome in dairy cattle using MPSS revealed strong similarities at functional level between rumen viral samples, which were highly distinct from the rumen microbiome samples. Significant variation between animals living in different housing while animals housed together presented similar viromes (Bathe et al., 2003). Ross and co-workers also observed large effects of feed additives on the rumen microbiome profiles using MPPS (Ross et al., 2013b). Recently, Earl et al., (2018), analysed the 16S rRNA microbiome surveys using massively parallel DNA sequencing technologies and reported an increased taxonomic and phylogenetic resolution with species-level classification of >90% of taxa and relative abundances microbial population have been reported. They concluded that application of MPSS to marker genes could help enhance taxonomic assignments of microbial species and reference databases and subsequently increase the specificity of relationship between microbial communities (Earl et al., 2018).

**Non-PCR based next generation sequencing technology**

With recent advancements in NGS, several PCR-independent NGS technologies have evolved which does not require prior DNA amplification using PCR. Some of these emerging sequencing technologies include single molecule sequencing with Heliscope single molecule sequencer, PacBio single molecule real-time (PacBio SMRT) sequencer and Oxford nanopore sequencer systems which need no pre-amplification steps. Pacific biosciences single molecule real-time (SMRT) sequencing technology is another molecular microscope with no requirement for template amplification step. It enables single molecule detection using sensitive optics which could spot individual fluorescently labeled nucleotide (Buermans and den Dunnen, 2014). These technologies are reliable, fast and sensitive enough to detect signal and read the DNA sequence of individual molecule template extensions. Thus, providing extension of un-amplified molecules with greater flexibility in the kinetics of the sequencing chemistry (Pushkarev et al., 2009; Zhou et al., 2013). Illumina shotgun sequencing (such as HiSeq, MiSeq), is another PCR-independent sequencing procedure
that enables estimation of species composition without PCR as a pre-step (Zhou et al., 2013).

**RFLP/T-RFLP and DGGE/TGGE**

Restriction fragment length polymorphism (RFLP), or terminal-restriction fragment length polymorphism (T-RFLP), (allows profiling, but also quantitation of microbiota). Denaturing gradient gel electrophoresis (DGGE), or temperature gradient gel electrophoresis (TGGE), are commonly used PCR based molecular fingerprinting techniques for evaluating microbial community and diversity. T-RFLP uses fluorescent-labelled primer for amplification of bacterial DNA fragments during PCR and this allows profiling as well as quantitation of microbial communities using capillary electrophoresis of PCR products fragmented by size with sequence-specific restriction enzymes. During DGGE analysis, a gel containing a linear gradient of DNA denaturants is used, while TGGE a temperature gradient is used for separation. DGGE/TGGE has shorter bands (20 to 40 bands) though many bacterial phylotypes will have similar or the same melting characteristics. This is a key factor that limits visualisation and evaluation of changes in only predominant bacterial groups. There are possibilities to resect and sequence individual band of interest. These techniques separate the PCR amplicons that were generated with either universal or group-specific primers to yield a profile (fingerprint) of the bacterial community (Suchodolski, 2012).

RFLP has been applied in combination with PCR procedures in comparing the genetic diversity and phylogenetic experiments. Avguštin et al. (1994) aimed to evaluate the nature of diversity that exist among 29 strains of *Prevotella ruminicola* from the rumen has identified signature oligonucleotides based on 16S ribosomal DNA sequences which distinguished the strains related to strains 23T, B(1)4, GA33, and M384, as well as an oligonucleotide that specifically recognised all but one of the *Bacteroides* and *Prevotella* strains tested. 5 ammonia hyper-producing (HAP) bacteria, using RFLP of 16S rDNA indicated that isolates differed from the previously described HAP bacteria and 16S rDNA PCR-RFLP-based investigations suggested that ruminal *Strptococcus bovis* diversity may be a diet-dependent phenomenon (Kleiβ et al., 1999; Jarvis et al., 2001). Regensbogenova et al., (2004b) reported that restriction fragment length polymorphism (RFLP) analysis of PCR amplified 18S rDNA sequences could be applied in identifying different rumen protozoa species. However, researchers were of the opinion that since 18S rDNA sequences among different protozoa are found to be closely related, the phylogenetic resolution of such RFLP analysis remains unclear (McSweeney et al., 2007).
The crucial element of RFLP as a procedure is the selection of an efficient and suitable restriction enzymes for the analysis and is applied in exploring the composition of individual bacterial species (Dang et al., 2007). Schlegel et al., (2003), used two strains of streptococcus, S. bovis and S. equinus which are important intestinal bacterial isolates from human and animal specimens. Schlegel et al. (2003), investigated the nature of the diversity of large bacterial complex including different species frequently isolated from infections of humans (Streptococcus galolyticus, Streptococcus infantarius) or animals (S. bovis, S. equinus, S. alactolyticus). The diversity of strains of S. bovis biotype II was analysed, and it was confirmed that they belong to different species, either S. equinus or S. infantarius. Tymensen et al. (2012), used T-RFLP to analyse rumen protozoa communities. The protozoa communities from hay-fed and silage/grain-fed cattle were different while multivariate analysis indicated that cattle fed the same diet and housed together (pen mates) tended to have similar protozoa community types.

Analysis of ruminal methanogenic Archaea communities in the rumen using DGGE is an efficient PCR-based technique in revealing the diversity and complexity of the rumen microbial community. It separates PCR products of the same length but differing in sequence since different sequences possess different melting temperatures resulting in contrasting migration behaviour. Additionally, analysis of PCR-DGGE fingerprint and phylogenetic analysis 16S rDNA sequences in DGGE profiles were combined to reveal the dominant bacterial communities and compared the variations of such bacterial communities in cattle breed. Leng et al. (2011) (consistent with Yang et al., 2010) shows that the dominant bacteria in the rumen phyla are Firmicutes, Bacteroidetes and Actinobacteria with significant variation in bacterial specie according to the cattle breeds investigated. Székely et al. (2009), compared the accuracy of DGGE and T-RFLP using bacterial diversity in mature compost bacterial community and cloning of environmental 16S rDNA and reported that DGGE shows less diversity (15–22 bands per sample) than T-RFLP (20–59 peaks greater than or equal to 1% of total peak area per sample). However, principal component analysis (PCA), of the DGGE and T-RFLP profiles showed that both molecular fingerprinting techniques gave a similar clustering of the samples.

**CONCLUSION**

The development and thorough application of PCR/qPCR DNA-based technologies in rumen microbiology research have provided a deeper insight into the biological understanding of the rumen microbiota. It is precise, more
PCR AND QPCR-BASED APPLICATIONS IN RUMEN MICROBIOLOGY RESEARCH

reliable and more efficient than traditional, culture-based identification and characterisation of the rumen microbiota. Through the use of PCR/qPCR-based procedures, quantitation of different rumen microbiota as well as analysis of phylogenetic relationship between and within these various microbial constituents in the rumen is achievable. However, it is advisable to optimise these procedures and other “omic”-based technologies used with PCR/qPCR protocols for accurate analysis of the complex rumen microbiome.

ACKNOWLEDGEMENTS

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ACTA AGRARIA KAPOSVÁRiensis

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