Symposium review: Uncertainties in enteric methane inventories, measurement techniques, and prediction models


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ABSTRACT

Ruminant production systems are important contributors to anthropogenic methane (CH$_4$) emissions, but there are large uncertainties in national and global livestock CH$_4$ inventories. Sources of uncertainty in enteric CH$_4$ emissions include animal inventories, feed dry matter intake (DMI), ingredient and chemical composition of the diets, and CH$_4$ emission factors. There is also significant uncertainty associated with enteric CH$_4$ measurements. The most widely used techniques are respiration chambers, the sulfur hexafluoride (SF$_6$) tracer technique, and the automated head-chamber system (GreenFeed; C-Lock Inc., Rapid City, SD). All 3 methods have been successfully used in a large number of experiments with dairy or beef cattle in various environmental conditions, although studies that compare techniques have reported inconsistent results. Although different types of models have been developed to predict enteric CH$_4$ emissions, relatively simple empirical (statistical) models have been commonly used for inventory purposes because of their broad applicability and ease of use compared with more detailed empirical and process-based mechanistic models. However, extant empirical models used to predict enteric CH$_4$ emissions suffer from narrow spatial focus, limited observations, and limitations of the statistical technique used. Therefore, prediction models must be developed from robust data sets that can only be generated through collaboration of scientists across the world. To achieve high prediction accuracy, these data sets should encompass a wide range of diets and production systems within regions and globally. Overall, enteric CH$_4$ prediction models are based on various animal or feed character-
istic inputs but are dominated by DMI in one form or another. As a result, accurate prediction of DMI is essential for accurate prediction of livestock CH4 emissions. Analysis of a large data set of individual dairy cattle data showed that simplified enteric CH4 prediction models based on DMI alone or DMI and limited feed- or animal-related inputs can predict average CH4 emission with a similar accuracy to more complex empirical models. These simplified models can be reliably used for emission inventory purposes.

**Key words:** enteric methane, uncertainty, prediction model, livestock

**INTRODUCTION**

The livestock sector is a significant source of anthropogenic greenhouse gas (GHG) emissions. In the United States, emissions from livestock production contributed an estimated 48% of the 2015 agricultural GHG emissions (US EPA, 2017). In Europe (EU-28), 59% of estimated agricultural GHG emissions were from livestock in 2015 (http://ec.europa.eu/eurostat/web/agriculture/data/database; accessed December 5, 2017). Methane (CH4) and nitrous oxide are the 2 most important GHG from agricultural activities. Methane, a potent short-lived (12.2-yr lifetime; Myhre et al., 2013) GHG, is emitted from livestock operations through enteric fermentation in the animal’s gastrointestinal tract (reticulo-rumen and hindgut) and similar methanogenic processes in manure. Globally, enteric CH4 emissions make up about one-fifth of the 10 to 12 Gt CO2-equivalent/yr GHG emissions from the Agriculture, Forestry, and Other Land Use sector (IPCC, 2014). There are, however, large uncertainties associated with estimating GHG emissions from livestock (or any other source), which has led to discrepancies between top-down (i.e., based on atmospheric measurements) and bottom-up (based on national or regional activity data and emission factors for different CH4 sources) and among bottom-up CH4 emission inventories (Miller et al., 2013; Hristov et al., 2014, 2017; Wecht et al., 2014; Maasakkers et al., 2016). These uncertainties may be related to uncertainties in changes in CH4 sinks (Rigby et al., 2017), or to uncertainties in changes in CH4 sources. As an example, a recent bottom-up inventory analysis, based mostly on national inventory reports, suggested that global livestock CH4 emissions are 11% greater than estimates based on Intergovernmental Panel on Climate Change (IPCC) emission factors (Wolf et al., 2017). As an 11% difference is well within the uncertainty bounds for livestock CH4 inventories (Hristov et al., 2017; US EPA, 2017), conclusions from such analyses have to be interpreted with caution. Therefore, the objective of this paper was to review uncertainties and discrepancies in CH4 inventories as related to livestock emissions, enteric CH4 measurement methods, and DMI and CH4 prediction models. The review and data presented here are an integral part of the GLOBAL NETWORK project and the Feed and Nutrition Network (http://animalscience.psu.edu/fmn/current-research/global-network-for-enteric-methane-mitigation; accessed December 4, 2017) within the Livestock Research Group of the Global Research Alliance for Agricultural Greenhouse Gases (www.globalresearchalliance.org; accessed December 4, 2017).

**UNCERTAINTIES IN ATMOSPHERIC METHANE CONCENTRATIONS AND ATTRIBUTION TO LIVESTOCK SOURCES**

Globally, atmospheric mixing ratio of CH4 (the number of moles of CH4 per mole of air) was relatively stable between 1999 and 2006 but have increased continuously since 2006 at a rate of 4 to 12 nmol/mol per year (https://www.esrl.noaa.gov/gmd/cegg/trends_ch4/#global_growth; accessed June 16, 2017). There is no consensus about the major drivers for this increase and, in addition, there is considerable disagreement regarding the contribution of livestock to global CH4 emissions. Reports based on isotopic composition of CH4 in the atmosphere, ice cores, and archived air, or combined data from bottom-up and top-down methodologies suggested that post-2006 increases in CH4 emissions are predominantly caused by increases in microbial CH4 (Nisbet et al., 2016; Saunois et al., 2016; Schaefer et al., 2016). Microbial, or biogenic, CH4 is generated by methanogenic archaea and can be from wetlands and agricultural activities, mainly livestock production and rice cultivation (Stolper et al., 2015). The atmospheric mixing ratio of CH4 is a function of emissions and sinks. The major sink for atmospheric CH4 is oxidation by hydroxyl radicals (OH), occurring mostly in the troposphere, which accounts for approximately 90% of the global CH4 sink (Kirschke et al., 2013). Because of the short lifetime of OH, direct observations of atmospheric OH mixing ratio are difficult to accomplish (Rigby et al., 2017). Therefore, the increase in atmospheric CH4 cannot be reliably attributed to an overall increase in emissions. The analysis by Rigby et al. (2017) pointed to “significant OH-related uncertainties” in the atmospheric CH4 budget and concluded that it is impossible to implicate global CH4 emission changes as the primary driver for recent trends in atmospheric CH4 mixing ratio.

If there was an increase in atmospheric CH4 mixing ratio and the increase was caused by agricultural sources, specifically livestock emissions, the trends in atmospheric CH4 should correspond to dynamics in global
livestock populations. During 1999 to 2006, however, when atmospheric CH\textsubscript{4} mixing ratio plateaued, global cattle and buffalo populations (these species make up 84% of all livestock enteric CH\textsubscript{4} emissions; FAOSTAT, 2017) continued to increase from 1.46 (1999) to 1.59 (2006) billion head (FAOSTAT, 2017), at a rate of approximately 18.8 million head/yr, which apparently did not affect atmospheric CH\textsubscript{4} over the same period. Since 2006, the rate of increase for the populations of these ruminant species declined to 7.3 million head/yr (FAOSTAT, 2017); we note that FAOSTAT does not specify uncertainty for their estimates, which is likely large for cattle inventories (and emission factors) in developing countries. Thus, it appears that the global dynamics in large ruminant inventories do not support the suggested farmed livestock origin of the increase in atmospheric CH\textsubscript{4} from 2006 to 2015. Potential increases in CH\textsubscript{4} emission from non-livestock agricultural sources to the global CH\textsubscript{4} budget cannot be excluded. Globally, the area harvested for paddy rice (emissions from which are typically 22 to 24% of the emissions from livestock), for example, had increased 42% from the 1960s to 2015 (FAOSTAT, 2017), although new rice varieties (i.e., water-saving and drought-resistance rice, or WDR; Luo, 2010) require less water and thus emit less CH\textsubscript{4} (Sun et al., 2016).

Source attribution of atmospheric CH\textsubscript{4} is largely based on its stable isotope signature, specifically \(^{13}\text{C}/^{12}\text{C}\). The average isotopic signature of microbial CH\textsubscript{4} appears to be quite distinct from that of fossil fuel CH\textsubscript{4} (Wang et al., 2015; Schwietzke et al., 2016). In the Wang et al. (2015) study, average \(\delta^{13}\text{C}\) of thermogenic CH\textsubscript{4} from the Northern Appalachian Basin was −36.2 to −25.7 ‰, whereas \(\delta^{13}\text{C}\) of enteric CH\textsubscript{4} from cows from the Pennsylvania State University’s dairy herd was −54.2 to −52.8 ‰. Based on CH\textsubscript{4} isotopic signature data, Schwietzke et al. (2016) concluded that fossil fuel CH\textsubscript{4} emissions are not increasing over time, implying that emissions of CH\textsubscript{4} from microbial sources have been increasing. Examination of the \(\delta^{13}\text{CH}_4\) database used in the Schwietzke et al. (2016) study (https://www.esrl.noaa.gov/gmd/ccgg/d13C-src-inv/; accessed December 4, 2017), however, shows a relatively large variability and uncertainty in the \(\delta^{13}\text{CH}_4\) data, from −68% (SD = 3.0%) for C3 plant–based ruminant diets to −54% (SD = 3.0%) for C4 plant diets; the authors used \(\delta^{13}\text{CH}_4\) of −66.8 ± 2.8% as a global average for ruminants, which is very close to that for wetlands (−61.5 ± 0.6%). Wang et al. (2015) also reported similar \(\delta^{13}\text{CH}_4\) for ruminal and swamp CH\textsubscript{4} samples. In the Schwietzke et al. (2016) database (over 8,100 observations), \(\delta^{13}\text{CH}_4\) of fossil fuel CH\textsubscript{4} (average of −45.0 ± 6.96‰ with minimum and maximum of −64.1 and −29.1‰, respectively) had a standard deviation as high as 15 to 16‰. This large variability in the isotopic signatures of microbial and fossil fuel CH\textsubscript{4} requires a more cautious interpretation of the data on CH\textsubscript{4} emission source distribution and the conclusions of Schwietzke et al. (2016). Furthermore, a recent analysis by Turner et al. (2017) showed significant overlap in the \(\delta^{13}\text{CH}_4\) isotopic signatures of fossil fuel (−15 to −76‰) and non-fossil-fuel (−31 to −93‰) CH\textsubscript{4} sources. As pointed out by Turner et al. (2017), fossil fuel CH\textsubscript{4} is not entirely thermogenic in origin (based on its isotopic signature), with over 20% of the world’s natural gas reserves generated by microbial activities (i.e., carrying biogenic isotopic signature). Thus, collectively, we can conclude that quantitative attribution of changes in atmospheric CH\textsubscript{4} concentrations to CH\textsubscript{4} sources based on \(\delta^{13}\text{CH}_4\) data is at least questionable. Both enteric and manure emissions contribute to livestock CH\textsubscript{4}, with manure reportedly being less depleted in \(^{13}\text{C}\) than enteric CH\textsubscript{4}, which further decreases the usefulness of the \(\delta^{13}\text{CH}_4\) signature approach for estimating the share of microbially derived CH\textsubscript{4} (Klevenhusen et al., 2010). Additional isotope measurements such as \(^{13}\text{CH}_4\), hydrogen isotopes, deuteromethane, or clumped isotopes (heavy isotopes that are bonded to other heavy isotopes; Eiler, 2007; Stolper et al., 2015; Wang et al., 2015) would help better discriminate individual source contributions.

**UNCERTAINTIES IN LIVESTOCK METHANE INVENTORIES**

Globally, estimated non-CO\textsubscript{2} GHG emissions from agriculture increased at a rate of 0.9%/yr between 1990 and 2010 (IPCC, 2014). In the United States, the Environmental Protection Agency (US EPA, 2017) reported a 16% decrease in CH\textsubscript{4} emissions between 1990 and 2015, due mainly to estimated decreases in emissions associated with fossil fuel exploration and production. The EPA’s bottom-up CH\textsubscript{4} inventory was challenged by top-down analyses suggesting that livestock CH\textsubscript{4} emissions are underestimated by as much as 80% by the EPA (Miller et al., 2013; Wecht et al., 2014). In the Wecht et al. (2014) study, oil and gas emissions, the largest source of anthropogenic CH\textsubscript{4} in the United States, were estimated to be 20% lower than EPA’s bottom-up estimates. A more recent top-down analysis indicated a sharp 30% increase in anthropogenic CH\textsubscript{4} emissions in the United States between 2002 and 2014 (Turner et al., 2016). According to their study, the spike in atmospheric CH\textsubscript{4} was mainly over the central part of the United States. Although the authors (Turner et al., 2016) mentioned a 20% increase in oil and gas production and a 9-fold increase in shale gas production in the United States (from 2002 to 2014), they concluded that the data do not allow attribution.
of atmospheric CH\textsubscript{4} mixing ratio to a specific source. It is worth pointing out that the cattle population (the major source of livestock enteric and manure CH\textsubscript{4} emissions) in the United States has been declining since the late 1970s, from 111 million in 1980 to 92 million in 2016 (NASS, 2017). Body weight of beef (and dairy) cattle has been increasing, however; as an example, despite the decreasing beef cattle numbers, total beef slaughter production has increased from about 107 to 125 million kilograms from 1980 to 2016 (NASS, 2017). This increase in the live and carcass weight of cattle, which likely corresponds to greater DMI, will partially offset the potential decrease in enteric CH\textsubscript{4} emission from the beef sector in the United States, caused by decreasing cattle inventories.

The uncertainties in livestock enteric CH\textsubscript{4} emissions in the current US EPA (2017) report are −11 and 18% (lower and upper bounds, respectively), corresponding to a 95% confidence interval, with the lower bound corresponding to the 2.5th percentile and the upper bound corresponding to 97.5th percentile, respectively. For CH\textsubscript{4} emissions from manure management, the uncertainty is −18 and 20%, respectively (US EPA, 2017). These uncertainties result from several factors, including uncertainties in animal inventories, DMI, ingredient and chemical composition of the diet, and CH\textsubscript{4} emission factors (for enteric fermentation) and inaccuracies of measurement of CH\textsubscript{4} emission from manure (minute amounts, often emitted as bubbles) related to manure composition, manure management system, duration of manure storage, and environmental factors such as temperature and wind. A recent gridded (0.1° × 0.1° grid; which represents an area of 81 to 109 km\textsuperscript{2}) inventory of livestock CH\textsubscript{4} emissions in the continental United States reported lower and upper 95% confidence bounds of −15.6 and 16.9% (as % of the mean; enteric), −65.0 and 63.3% (manure), and −19.3 and 19.2% (total emissions), respectively (Hristov et al., 2017). In that analysis, major sources of uncertainties for enteric CH\textsubscript{4} were animal BW (lower and upper 95% confidence bounds across cattle categories: −18 to −24% and 21 to 29%, respectively), DMI (−21 to −29% and 21 to 29%), and CH\textsubscript{4} yield (−18 to −41% and 19 to 42%). In a model designed to estimate enteric CH\textsubscript{4} from Dutch dairy farms, Bannink et al. (2011) reported that the largest uncertainty (18%) was related to VFA stoichiometry. Estimates for total livestock CH\textsubscript{4} emissions in the Hristov et al. (2017) study were comparable to current US EPA (2017) estimates for 2012 (last census of agriculture) and to estimates from the gridded Emission Database for Global Atmospheric Research (EDGAR, 2011) inventory. However, the spatial distribution of emissions in the Hristov et al. (2017) analysis differed significantly from that of EDGAR and a recent gridded inventory based on US EPA’s emission database (Maasakkers et al., 2016). For example, the combined enteric and manure CH\textsubscript{4} emissions from livestock in Texas and California (the largest contributors to the national total) in the Hristov et al. (2017) study were 36% lower and 100% greater, respectively, than estimates from EDGAR. These differences originate from differences in emission factors between the 2 analyses [lower emission factors for feedlot cattle (i.e., Texas) and higher emission factors for dairy cows (i.e., California) in the Hristov et al., 2017 analysis]. Gridded bottom-up emission inventories, such as EDGAR, are commonly used to assess the contribution of CH\textsubscript{4} from different sectors within a region. Top-down approaches use these bottom-up inventories as a prior estimate of total emissions and, in some cases, to allocate the resulting (posterior) emission estimates to emission sources (Saunois et al., 2016). As a result, spatial distribution of emissions in gridded inventories likely strongly affects the conclusions of top-down approaches that use them, especially in the source attribution of emissions (i.e., biogenic vs. thermogenic or livestock vs. fossil fuel); therefore, conclusions from such studies should be interpreted with caution, even more when aiming to make future projections and evaluate mitigation options.

**UNCERTAINTIES IN ENTERIC METHANE MEASUREMENT TECHNIQUES**

Several established techniques exist for direct measurement of enteric CH\textsubscript{4} emissions from ruminants. These include respiration chambers (RC), the sulfur hexafluoride (SF\textsubscript{6}) tracer technique, and more recently, the GreenFeed technique (GF; C-Lock Inc., Rapid City, SD), which is an automated head-chamber system. In addition, several indirect techniques have also been proposed and used for measuring enteric CH\textsubscript{4} emissions (reviewed by Negussie et al., 2017). A comprehensive review of current enteric CH\textsubscript{4} measurement techniques was recently published by an international team of scientists (Hammond et al., 2016a) as part of the GLOBAL NETWORK project.

The GLOBAL NETWORK project has collected thousands of measurements of CH\textsubscript{4} emissions from individual animals and accompanying data (e.g., diet composition and DMI) to develop robust, broadly applicable CH\textsubscript{4} prediction equations for applications such as livestock CH\textsubscript{4} inventories. Contributors supplying data to the GLOBAL NETWORK project used various methods for measuring enteric CH\textsubscript{4}. Three databases were created, one each for dairy cows, beef cattle, and small ruminants (sheep and goats). In Table 1, we present data for the main measurement techniques that
were included in the dairy database of the GLOBAL NETWORK project. The RC sub-database included cows with DMI and milk yield that were lower than those of cows included in the GF sub-data set but comparable to those in the SF6 data set. Also, the range of DMI was narrower for GF and SF6 than for RC. As evident from the data, significant variation was associated with all measurement methods for CH4 emission rate, yield, and intensity; the coefficient of variation (CV) for emission rate (g of CH4/d) averaged 30, 18, and 28% for RC, GF, and SF6, respectively. It is important to note that the variability included in these CV values includes all sources of variation, not just variation due to method of measurement and how it was used. Methane emission rate is determined primarily by the amount of rumen fermentable substrate and, for this reason, comparisons of CV are better made based on CH4 yield; that is, grams of CH4 per kilogram of DMI. On this basis, the CV for RC is reduced to 21% and is comparable to that for GF and SF6 (21 and 27%, respectively). Low variability, however, does not always mean high accuracy. Each method has to be carefully evaluated by researchers who, based on their expertise and available data, can determine whether a method can be reliably used to measure enteric CH4 emission from ruminants for the specific conditions and objectives of their experiment and animals used.

**Respiration Chambers**

Respiration chambers have been considered the gold standard for measuring enteric CH4 emission from farm animals, although this is only the case if RC are operated properly and recoveries are fixed and preferably close to 100%. Moreover, there are many kinds of chambers and operation procedures with varying accuracies. As shown in a collaborative project in the United Kingdom, RC can also produce inaccurate results (Gardiner et al., 2015). In that ring-test, measured CH4 recovery was unacceptably low for several of the RC tested. Critical sources of variation for measurement of CH4 emission through RC are airflow rate through the chamber and the dynamics of air mixing in the chamber, which determines response time. In the ring-test by Gardiner et al. (2015), 3 potential sources of experimental error were evaluated by testing the measured recovery of a reference source of ultra-high-purity CH4 standard released at calibrated rates at specific points in the chambers to test the accuracy of specific components of the measurement system. The tested sources of error were analyzer error, ducting efficiency (from chambers to analyzers, including measurements of airflow), and mixing of air in chamber. Of these, ducting and airflow measurement were the largest source of variation in CH4 standard recovery within and between RC and research facilities (1.3, 15.3, and 3.4% variation for analyzers, ducting/flow, and air mixing in chamber, respectively). Chambers need to be routinely calibrated and demonstrate gas recovery rates of approximately 100% both before and after each experimental deployment, as highlighted recently by Gerrits et al. (2018).

As well as these issues, several other common but often overlooked issues can influence CH4 yield measurements made using RC. Animals in RC must have stable daily feed intake. Moate et al. (2012) showed that, for a dairy cow in RC, approximately 30% of today’s CH4 emissions are a result of yesterday’s DMI. It is commonly observed that dairy cows may slightly reduce their DMI on the first day they enter a respiration chamber (data from the first day are normally excluded from the analysis). Thus, day-to-day variation in total DMI can cause an error in estimated CH4 yield of up to 3% (Moate et al., 2012). If RC are fitted with air locks for entry and feeding, disruption to measurements is minimized, the entry and presence of staff in the RC can be accounted for (see Reynolds and Tyrrell, 2000), and measurements can be obtained without interruption for successive 24-h periods (Flatt et al., 1958; Tyrrell et al., 1979). However, many modern RC are constructed such that the chamber doors must be opened for approximately 30 min at least twice per day to enable milking and cleaning. With exclusion of these time slots, CH4 measurements from a specific chamber may cover approximately 23 h/d. There does not appear to be an internationally agreed protocol for filling the total 1-h “gap” in missing CH4 measurements. Interpolation may be used for this purpose but what approximation should be used for the missing data? This would not be a problem if the rate of CH4 emissions were constant over the course of a day, but with dairy cows, there is often considerable hour-to-hour variation in rate of CH4 production, with the peak hourly rate of CH4 emission being more than 3 times the minimum hourly rate of CH4 emission. Depending on feeding (immediately upon entrance or just before leaving the chamber), the most accurate estimate of CH4 production rates during the two 30-min gap periods is the average of the CH4 production immediately preceding and after each opening, or the CH4 production rate immediately preceding each opening of the chamber. However, the most common practice is to use the mean rate of CH4 production as measured during the 23 h for which data are available. The latter interpolation method can result in an overestimation of CH4 emission and hence CH4 yield by approximately 2% (P. J. Moate, unpublished data). In contrast, van Gastelen et al. (2017) established a very small difference of 0.1% in daily CH4 emission rate when comparing discarding
Table 1. Descriptive statistics of enteric methane emission, measured using direct methods, DMI, and milk and 3.5% fat- and protein-corrected milk yields used in the analysis (data from the GLOBAL NETWORK project; Niu et al., 2018)

<table>
<thead>
<tr>
<th>Method</th>
<th>Geographic location and contributing laboratories</th>
<th>Variable</th>
<th>n</th>
<th>Mean</th>
<th>SD</th>
<th>Minimum</th>
<th>Maximum</th>
<th>CV</th>
<th>95% Confidence limits for mean</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>All data Europe, North and South America, Australia, and New Zealand (20 laboratories)</td>
<td>CH₄</td>
<td>4,152</td>
<td>357.8</td>
<td>104.6</td>
<td>67.8</td>
<td>728.6</td>
<td>29.2</td>
<td>354.6 361.0</td>
</tr>
<tr>
<td></td>
<td>CH₄DMI</td>
<td>4,152</td>
<td>20.1</td>
<td>4.3</td>
<td>4.4</td>
<td>38.1</td>
<td>21.6</td>
<td>19.9 20.2</td>
<td></td>
</tr>
<tr>
<td></td>
<td>CH₄/MY</td>
<td>3,983</td>
<td>15.6</td>
<td>8.5</td>
<td>2.3</td>
<td>119.6</td>
<td>54.2</td>
<td>15.4 15.9</td>
<td></td>
</tr>
<tr>
<td></td>
<td>DMI</td>
<td>4,152</td>
<td>18.1</td>
<td>4.8</td>
<td>3.9</td>
<td>35.4</td>
<td>26.4</td>
<td>17.9 18.2</td>
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</tr>
<tr>
<td></td>
<td>MY</td>
<td>3,983</td>
<td>26.7</td>
<td>10.5</td>
<td>1.3</td>
<td>62.7</td>
<td>39.2</td>
<td>26.4 27.1</td>
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</tr>
<tr>
<td></td>
<td>FPCM</td>
<td>3,865</td>
<td>28.4</td>
<td>10.6</td>
<td>1.5</td>
<td>65.4</td>
<td>37.2</td>
<td>28.1 28.7</td>
<td></td>
</tr>
<tr>
<td>RC</td>
<td>Europe, North America, Australia, and New Zealand (13 laboratories)</td>
<td>CH₄</td>
<td>3,024</td>
<td>344.5</td>
<td>103.2</td>
<td>67.8</td>
<td>701.0</td>
<td>30.0</td>
<td>340.8 348.2</td>
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<tr>
<td></td>
<td>CH₄DMI</td>
<td>3,024</td>
<td>20.2</td>
<td>4.2</td>
<td>4.4</td>
<td>36.1</td>
<td>20.9</td>
<td>20.0 20.3</td>
<td></td>
</tr>
<tr>
<td></td>
<td>CH₄/MY</td>
<td>2,874</td>
<td>16.1</td>
<td>9.1</td>
<td>2.3</td>
<td>119.6</td>
<td>56.8</td>
<td>15.8 16.4</td>
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</tr>
<tr>
<td></td>
<td>DMI</td>
<td>3,024</td>
<td>17.2</td>
<td>4.4</td>
<td>3.9</td>
<td>33.5</td>
<td>25.7</td>
<td>17.1 17.4</td>
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<tr>
<td></td>
<td>MY</td>
<td>2,874</td>
<td>25.4</td>
<td>10.2</td>
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<td>59.7</td>
<td>40.4</td>
<td>25.0 25.7</td>
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<td>57.6</td>
<td>39.0</td>
<td>26.5 27.3</td>
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<tr>
<td>GF</td>
<td>Europe, North America (4 laboratories)</td>
<td>CH₄</td>
<td>731</td>
<td>435.3</td>
<td>78.6</td>
<td>139.0</td>
<td>728.6</td>
<td>18.0</td>
<td>429.5 441.0</td>
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<tr>
<td></td>
<td>CH₄DMI</td>
<td>731</td>
<td>20.0</td>
<td>4.3</td>
<td>6.2</td>
<td>32.8</td>
<td>21.4</td>
<td>19.7 20.3</td>
<td></td>
</tr>
<tr>
<td></td>
<td>CH₄/MY</td>
<td>729</td>
<td>14.3</td>
<td>5.4</td>
<td>3.1</td>
<td>51.6</td>
<td>38.0</td>
<td>13.9 14.7</td>
<td></td>
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<tr>
<td></td>
<td>DMI</td>
<td>731</td>
<td>22.3</td>
<td>4.1</td>
<td>13.9</td>
<td>35.4</td>
<td>18.2</td>
<td>22.0 22.6</td>
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<td>65.4</td>
<td>24.5</td>
<td>34.9 36.2</td>
<td></td>
</tr>
<tr>
<td>SF₆</td>
<td>Europe, North and South America (6 laboratories)</td>
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1RC = respiration chambers; GF = GreenFeed system (C-Lock Inc., Rapid City, SD); SF₆ = sulfur hexafluoride tracer technique.
2CH₄ = methane emission, g/head per day; CH₄DMI = g of methane emission per kg of feed DMI (emission yield); CH₄/MY = g of methane emission per kg of milk yield (emission intensity); DMI = dry matter intake, kg/d; MY = milk yield, kg/d; FPCM = fat- and protein-corrected milk yield, kg/d (from Leiva et al., 2000, based on Tyrrell and Reid, 1965).
3n = number of observations in the data set.
(and interpolating between last time point before opening and first time point after closing the chamber) with not discarding the data from these time slots.

**The SF₆ Technique**

Another widely used technique to measure enteric CH₄ emissions is the SF₆ tracer method (Zimmerman, 1993; Johnson et al., 1994). Variability with the SF₆ technique has been notoriously high (Pinares-Patiño and Clark, 2008; Pinares-Patiño et al., 2011), but the modifications by Deighton et al. (2014) addressed the most important sources of error, and the modified technique produced CH₄ measurements with accuracy similar to measurements using RC. Part of the variation with SF₆ seems intrinsic to the technique because the estimated CH₄ emission rate appears sensitive to factors that affect the proportions of exhaled and eructated air in the air samples collected and distance of the sampling point from to the mouth/nostrils (Berends et al., 2014), which is not an issue with RC. Several important conditions must be met to reduce variability in the CH₄ measurement data when the SF₆ technique is used. These include (1) high and known release rate of SF₆ from the permeation tube, (2) at least 5 (depending on day-to-day variation in emission rates; Arbre et al., 2016) consecutive measurement days, and (3) low concentrations of SF₆ and CH₄ in the background air (i.e., using the technique in enclosed barns is not recommended, unless there is adequate ventilation throughout the measurement period; Dorich et al., 2015; Hristov et al., 2016). Even with adequate ventilation, samples of background air concentrations should always be included to correct the measurements obtained. In this regard, the method of obtaining background concentrations is important and should be as representative as possible of the background air in which the measurements are being obtained. A suitable approach is to include animals in the trial that are sampled in the same way as the other animals in the study but are not given an SF₆ permeation tube. Other concerns addressed by the studies of Deighton et al. (2014) include variation in release rate of permeation tubes over time (months) after calibration and variation in sampling rate over time (hours) during the sampling day, both of which can introduce bias in estimates obtained. Variation in release rate can be accounted for in part by using Michaelis-Menten kinetics to estimate the decay in release rate over time, rather than first-order kinetics (Deighton et al., 2014) if measurements are obtained more than 60 d after calibration of permeation tubes. Deighton et al. (2014) also showed that bias due to variation in sampling rate over the course of a 24-h sampling period is markedly reduced when orifice plate flow controllers, rather than capillary tubes, are used to obtain air samples. Because of diurnal changes in CH₄ emission over the course of each day, sampling for less than 24 h is not appropriate for estimates of daily rate of CH₄ emission. When these conditions and considerations are addressed, the SF₆ tracer technique can produce accurate CH₄ emission data from a large group of animals. In a review of CH₄ emission techniques, Hammond et al. (2016a) reported that, in 5 studies comparing CH₄ emissions from dairy cows obtained using RC and SF₆ (simultaneously in 2 studies), measurements of CH₄ emission were not significantly different in 4 studies and were different in 1 study (422 vs. 469 g/d). Detailed guidelines for using the SF₆ technique were published by an international panel of experts (Berndt et al., 2014).

**The GreenFeed System**

A more recent technique for direct measurement of enteric CH₄ emissions is the automated head-chamber system GreenFeed, which was developed for spot sampling of exhaled and eructated gases (Zimmerman and Zimmerman, 2012). When properly used (Hristov et al., 2015a), GF can be a reliable technique for measuring enteric CH₄ emissions from ruminant animals (Dorich et al., 2015; Hammond et al., 2016a,b; Hristov et al., 2016). An important prerequisite for decreasing uncertainty of the measurement when using GF is that all animals visit the unit at times that enable estimation of the diurnal pattern of CH₄ emission over successive 24-h periods. Methane emissions have a clear diurnal pattern related to the pattern of feed intake (usually lower at night; Brask et al., 2015; Hammond et al., 2016a); therefore, for accurate daily emission estimates, animal visits need to be distributed appropriately over the 24-h feeding cycle. The number and timing of visits to GF will vary depending on the type of animal, the diet fed, and the level of DMI (Hammond et al., 2016a,b). Reliable results with GF can be obtained when the number and timing of animal visits are controlled by the investigator, which is easily achievable in a tiestall barn situation (Branco et al., 2015; Hristov et al., 2015b; Dittmann et al., 2016). Alternatively, measurements have to take place over a prolonged period (up to 3 to 5 wk, depending on the study objectives; Arbre et al., 2016; Renand and Maupetit, 2016; Arthur et al., 2017). Obtaining measurements at specific time points from each animal on a study over a series of days increases precision and, as a result, can provide an accurate determination of treatment effects on CH₄ emission. However, the measurements obtained are not necessarily accurate estimates of daily emission rate, if the timing of measurements does not adequately account for the diurnal pattern of emission (Doreau
et al., 2018). For studies in which groups of animals are provided access to a GF unit (or units), timing of use can be influenced by programming the unit to only provide feed to animals at specific intervals, which encourages the animals to visit the unit at varied times throughout successive days. Nevertheless, in practice, the number of visits tends to be higher at specific times of the day (e.g., Hammond et al., 2015, 2016a,b) and may be influenced by the type of diet fed.

A recent evaluation of a large number of estimates of CH$_4$ emission rate (g/d) from 2 studies in growing beef cattle (Arthur et al., 2017) examined the number of observations (spot measurements) required to reliably estimate daily emission rate using GF, based on the reduction in variance observed with increasing number of observations. The authors found that as long as measurements were of sufficient duration (at least 3 min), 30 observations were sufficient to obtain reliable CH$_4$ emission data, regardless of how many times per day the measurements were obtained (on average 4.4 per day in one study and 1.3 per day in another), although the problem of unbalanced spread of visits over a 24-h period in view of diurnal CH$_4$ production patterns is not necessarily solved. These results emphasize the need for sufficient numbers of GF measurements per experimental unit (animal on a given treatment) for studies where animals are allowed voluntary access to the equipment. Another potential source of error in outdoor use is the effect of wind on the capture efficiency of the GF unit, which is used in the calculation of CH$_4$ emission rate for each measurement. Variation in wind speed and direction can affect measurements (Huhtanen et al., 2015a); thus, it is recommended that units used outdoors be fitted with anemometers to record wind speed during measurements so attempts can be made to correct measurements for the effects of wind. Measurements obtained using GF, similar to those obtained using the SF$_6$ technique, do not include CH$_4$ emissions from the rectum, but these emissions are typically small (approximately 1–3%, as measured or estimated by Murray et al., 1976 and Muñoz et al., 2012, respectively).

Overall, both GF and SF$_6$ are established techniques and can produce accurate estimates for enteric CH$_4$ emission when properly used and calibrated. Emphasis on further improvement of the methodology and experimental set-up (Deighton et al., 2014; Hristov et al., 2015a) will increase the accuracy of these techniques. Direct comparisons of GF and SF$_6$ with RC have shown acceptable agreement in some studies (e.g., Grainger et al., 2007; Muñoz et al., 2012; Deighton et al., 2014; Hammond et al., 2016b; Velazco et al., 2016; Jonker et al., 2016; Huhtanen et al., 2018; Alemu et al., 2017; Rischewski et al., 2017) but not in others (e.g., Pinares-Patiño et al., 2011; Hammond et al., 2015). The modified SF$_6$ technique, as proposed by Deighton et al. (2014), showed good agreement with RC; CH$_4$ yield was not different between SF$_6$ and RC, and the between-animal CV were similar between the 2 techniques (6.5 and 7.5%, respectively). A recent meta-analysis showed a strong relationship ($R^2 = 0.92$) between CH$_4$ emissions measured in RC and by GF used in the same experiment (Figure 1; Huhtanen et al., 2018). Sources of uncertainties with both techniques have been discussed above. To reduce variability in data generated by SF$_6$ or GF, researchers have to strictly follow recommended procedures or adjust these procedures to their specific experimental conditions when necessary.

**Indirect Methods**

Indirect approaches have been proposed and used to measure enteric CH$_4$ emissions in livestock. Usually, these methods are associated with lower accuracy and greater uncertainty in the emission data than the direct methods described above. One approach used estimated CO$_2$ emission and measured CO$_2$:CH$_4$ ratio in exhaled air to estimate CH$_4$ emission (Madsen et al., 2010). Changes in digestive and metabolic activities (even at the same level of feed intake), differences in feed efficiency, as well as variation in ruminal fermentation can all influence the amount of CO$_2$ produced by the animal and thus affect the predicted CH$_4$ emission (Huhtanen et al., 2015a). The CO$_2$:CH$_4$ ratio technique is comparable to the SF$_6$ technique in some ways, but
it is usually based on “spot” measurements of breath CH₄ concentration, rather than integrated measurements over 24 h, and the emission rate of the “tracer” gas (CO₂) is estimated, rather than relying on emission from a calibrated delivery device in the rumen, as with the SF₆ technique. Haque et al. (2017) evaluated CH₄ production calculated using observed CO₂ production in RC versus using CO₂ production calculated based on the heat production method of Madsen et al. (2010). In that evaluation, CH₄ production estimated using calculated CO₂ production resulted in smaller differences and changed the significance of treatment effects between diets compared with using the actual observed CO₂ production.

Another indirect method proposed by Garnsworthy et al. (2012) relies on estimating CH₄ emission during an eructation event and the frequency of eructation during a measurement period—the “sniffer” method. A feature of the method is that hundreds of repeated measurements can be made at little additional cost over prolonged periods. In 2 experiments with lactating cows, however, Huhtanen et al. (2015a) found larger variability with the sniffer method and no relationship to emissions measured using GF. Distance from the sampling inlet had a strong influence on measured gas concentration in a laboratory study and, in an animal study, the measured CH₄ concentration was strongly related to head position (Huhtanen et al., 2015a). In addition, head position was a highly repeatable characteristic precluding that an increased number of observations could solve the problem. Another recent study concluded that the capability of the sniffer method to adequately measure and rank CH₄ emission rates among dairy cows is highly uncertain and requires further investigation into the sources of variation (Wu et al., 2018).

Another indirect technique uses a laser CH₄ detector to measure CH₄ mixing ratio in the air between the laser device and the animal (usually 1 to 3 m). The method allows CH₄ measurements in on-farm conditions and from a large number of animals; however, comparative studies found a positive but weak relationship between the laser method and RC measurements (Chagunda et al., 2013; Ricci et al., 2014), although the device was found to accurately record variations in CH₄ in spent air of RC (Sorg et al., 2017). Environmental factors such as temperature, wind velocity (particularly important for grazing conditions), proximity of other animals, humidity, and others can affect the accuracy of the measurements. Further critical evaluation of these indirect methods has been provided in Hammond et al. (2016a), but as the methods are “indirect,” they rely on assumed relationships between concentrations of CH₄ in breath and other parameters and as such are subject to greater variance and uncertainty than direct measures of CH₄ emission rate.

**UNCERTAINTIES IN PREDICTING ENTERIC METHANE EMISSIONS**

**Relationship of DMI with CH₄ Emission and Prediction of DMI**

Dry matter intake is an important factor in enteric CH₄ prediction models. Models predicting DMI can be used in conjunction with emission factors to estimate enteric CH₄ emissions in a Tier 2 approach (which is based on country-specific emission factors and other data). Appuhamy et al. (2016) evaluated 40 prediction equations using data that included measured DMI and feed quality attributes. The best performing models in each region (North America, Europe, and Australia and New Zealand) were then re-evaluated using predicted DMI and compared with estimates that used measured DMI. Appuhamy et al. (2016) reported that models using estimated DMI predicted enteric CH₄ emissions as accurately as the measured data if DMI could be estimated with reasonable accuracy. Thus, enteric CH₄ emissions could be predicted well without DMI measurements for North America. For Europe, using estimated DMI rather than observed DMI resulted in satisfactory CH₄ emissions prediction. For Australia and New Zealand, CH₄ emissions could not be estimated well without actual DMI measurements. These differences were likely due to the models used. The DMI prediction model was developed based on North American data and may not work well with diets that have greater forage proportion, including cattle on pasture. In the GLOBAL NETWORK database of individual dairy cow data (Niu et al., 2018), CH₄ prediction equations with a greater number of independent variables performed best and had lower root mean squared prediction error (RMSPE) as a percentage of the mean observed value (14.7 to 19.8%). However, less complex models requiring only DMI had predictive ability comparable to those of the more complex models (RMSPE = 15.2 to 21.4%). This indicates that DMI alone may be sufficient to predict enteric CH₄ emissions for inventory purposes (as discussed in Hristov et al., 2017). The coefficient of determination for the relationship of measured CH₄ emissions with DMI, however, can be highly variable and may be influenced by several factors, including CH₄ measurement technique.

The relationships of measured CH₄ production and DMI (absolute or expressed on a BW basis) and NDF intake (NDFI) in the GLOBAL NETWORK dairy database (Niu et al., 2018) were investigated using the MIXED and REG procedures of SAS (version 9.4; SAS
Institute Inc., Cary, NC). Table 2 summarizes the results of these analyses. The linear relationship of DMI and CH₄ production was moderately strong (R² = 0.58) for the RC data (Figure 2, RC) and similar to the relationship for the entire data set (R² = 0.63; Figure 2, all data) but was very weak for GF (R² = 0.05; Figure 2, GF) and low for the SF₆ technique (R² = 0.27; Figure 2, SF₆); nonlinear models did not improve the relationship (data not shown). The estimated slopes indicate a much larger incremental yield in CH₄ with increasing DMI for RC than for GF and SF₆ (16.12 ± 0.299, 7.53 ± 0.775, and 5.87 ± 1.373 g of CH₄/kg of DMI, respectively). The prediction error was also lower for RC than for GF or SF₆. Similarly, relationships between DMI as a fraction of BW, NDFI, or milk yield or ECM yield and CH₄ were stronger for RC data than for GF or SF₆. This can be partially explained by the wider range of DMI data in the RC subset compared with that of GF or SF₆. The relationship of CH₄ emissions and DMI is usually strong with wider ranges of DMI (Hristov et al., 2013; Charmley et al., 2016) and weak when the range of DMI is narrower (Hristov et al., 2015b). The meta-analysis by Charmley et al. (2016) was on a large Australian data set (1,033 observations) including both dairy and beef cattle data and clearly showed that relationship between DMI and CH₄ emissions was strong (R² = 0.92) and the intercept was close to zero when DMI range was large (from about 2 to 28 kg/d in their analysis). If RC data in the current analysis were restricted to DMI >15 kg/d, R² for the relationship with DMI decreased to 0.41 and root mean squared error increased to 68.2 (data not shown).

A moderate relationship between DMI and CH₄ emissions has been established for both GF and SF₆ techniques. In a meta-analysis of dairy cow studies by Grainger et al. (2007), the relationship between DMI and CH₄ emission as measured by the SF₆ technique was R² = 0.56 and was better than the relationship between DMI and CH₄ emission for RC (R² = 0.39). The authors noted that in only 22% of the studies was the

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1All data = all data in the GLOBAL NETWORK project dairy data set; RC = data from studies using respiration chambers only; GF = data from studies using the GreenFeed system (C-Lock Inc., Rapid City, SD) only; SF₆ = data from studies using the sulfur hexafluoride tracer technique only.
2DMI = dry matter intake, kg/d; NDFI = neutral-detergent fiber intake, kg/d; BW = body weight, kg; DMI (or NDFI)/BW = DMI or NDFI as % of BW; MY = milk yield, kg/d; FPCM | 3.5% fat- and protein-corrected milk yield, kg/d (from Leiva et al., 2000, based on Tyrrell and Reid, 1965).
3n = number of observations in the data set.
4Mixed regression model analysis; all P-values <0.001.
5REG = fit statistics from a fixed regression model; RMSE = root mean squared error.
DMI of the cows >20 kg/d; more data are needed to establish a reliable relationship for greater DMI. A moderately strong relationship ($R^2 = 0.44$) of DMI and $CH_4$ emissions was demonstrated for GF in a beef data set (445 observations; DMI ranged from 3.6 to 19.1 kg/d) by Bird-Gardiner et al. (2017). In an experiment with dairy cows consuming around 28 kg of DM/d, however, the relationship of DMI with $CH_4$ emissions measured with GF or the $SF_6$ technique was relatively weak: $R^2 = 0.47$ and 0.08, respectively (Hristov et al., 2015b). The absence of a strong relationship between DMI and $CH_4$ emissions observed in the current analysis for both GF and $SF_6$, compared with the relationship for RC (Table 2 and Figure 2), is difficult to explain but reflects, in part, the variation associated with implementation the former techniques, as discussed earlier.

Most models developed to predict enteric $CH_4$ emissions usually include either DMI or some form of feed/nutrient intake; therefore, as pointed out earlier, accurate prediction of DMI is important for accurate prediction of $CH_4$ emissions and yield. The current dairy NRC (2001) model predicts DMI based on the cow’s metabolic BW, FCM yield, and stage of lactation. Dry matter intake prediction models for other categories of dairy cattle or beef cattle involve a variable for BW (metabolic BW or initial shrunk BW) and NE$_M$ concentration (NRC 2000, 2001, 2016). Numerous DMI prediction models have been proposed and evaluated (Ingversen, 1994; Mertens, 1995). An in-depth review of these models is outside the scope of this analysis and the examples given here are to illustrate the variable approaches (e.g., feed composition; animal factors...
such as BW, parity, and lactation stage; physiological mechanisms; genomic prediction of DMI. Understanding the factors important in regulating DMI in dairy cows.

Although it is generally agreed that DMI is the most important factor influencing CH$_4$ production, the general nature of this relationship remains undetermined. In the original equation proposed by Blaxter and Clapperton (1965), the relationship was curvilinear based on feeding level. More recently, Knapp et al. (2014) also proposed a curvilinear relation between DMI and CH$_4$, production, with CH$_4$ yield decreasing at high DMI. In dairy cows, very high DMI is usually only achieved with diets containing a relatively high proportion of concentrate feeds, and high concentrate diets are known to decrease CH$_4$ production (Blaxter and Clapperton, 1965). When the diet of cattle contains less than 30% concentrate, the relationship between DMI and CH$_4$ production has been shown to be linear, even to intakes up to 27 kg of DM/d (Charmley et al., 2016). A meta-analysis by Hristov et al. (2004) indicated that dietary concentrations of protein and carbohydrate fractions were important variables in predicting DMI in lactating dairy cows (and DMI was the dominant factor for estimating milk and milk protein yield). Shah and Murphy (2006) proposed an exponential DMI model based on lactation asymptotic maximum DMI and DIM. Zom et al. (2012) proposed a DMI prediction model based on estimated (from parity number, DIM, and days pregnant) feed intake capacity and a feed-specific satiety value, based on feed chemical composition and digestibility. The latter model and 4 other models (NRC, 2001 and 3 European models) were evaluated by Jensen et al. (2015). The models predicted DMI with various accuracies (RMSPE of 1.2 to 3.2 kg/d); best prediction was by a complex model involving BW, parity, DIM, milk yield, and dietary (forage) NE$_L$. An analysis of DMI prediction by 5 feeding systems yielded prediction errors of 1.6 to 3.2 kg/d (Krizsan et al., 2014). Appuhamy et al. (2018) evaluated the comprehensive (IPCC-CMP) and simplified (IPCC-SMP) IPCC models (IPCC, 2006), the modified Cornell Net Carbohydrate and Protein System model (CNCPS; Fox et al., 1992 as modified by Arnerdal, 2005), and the NRC (2001) models to predict DMI using an independent data set. The modified CNCPS, relying on BW and FCM yield, more accurately predicted DMI (RMSPE = 14.1%) than the NRC (RMSPE = 19.4%), IPCC-SMP (RMSPE = 16.9%), or IPCC-CMP (RMSPE = 23.4%) models. Overall, the results by Appuhamy et al. (2018) demonstrated that DMI can be predicted successfully using information such as milk yield and milk fat content (routinely available on dairy farms), which could therefore be used to estimate enteric CH$_4$ emissions.

**Prediction of CH$_4$ Emissions**

Prediction models have been widely used to estimate variation in CH$_4$ emissions for a variety of purposes (Kebreab et al., 2006). Many countries and regions of the world have set targets for the reduction of GHG emissions including CH$_4$. For example, California recently passed legislation mandating a reduction in the statewide emission of CH$_4$ by 40% below the 2013 levels by 2030 (State of California, 2017). Assessment of baseline emission in 2013 was determined using mathematical models, particularly those recommended by the IPCC (2006) and used in almost all national inventory protocols. Therefore, the accuracy of the model used is important in setting and assessing achievable targets. As existing models are based on limited databases, new and more-accurate models are required to establish the baseline for assessing any reduction in emissions or estimating global CH$_4$ emissions attributable to enteric fermentation. Where data sets used for CH$_4$ emission prediction model development are composed of data from multiple sources (e.g., different research groups and multiple studies) such as, for example, the GLOBAL NETWORK project, the effect of both research groups and studies should be incorporated in the model (Niu et al., 2018). In addition, if more than one CH$_4$ measurement technique was used by the same research group, the within-group variation from different techniques should also be considered.

**Types of Models Used to Predict Enteric CH$_4$ Emissions**

Enteric CH$_4$ emission predictions are obtained using different types of models. These range from simple emission factors (e.g., IPCC, 2006; Tier 1) and empirical models (e.g., Ramin and Huhtanen, 2013) to more detailed mechanistic models (e.g., Baldwin, 1995; Mills et al., 2001). Some models have been developed specifically to predict enteric CH$_4$ emissions from feed intake and other diet attributes (such as, for example, NDF and ether extract concentrations; e.g., Moraes et al., 2014); others have been modified or adapted to calculate emissions from ruminal fermentation kinetics (e.g., Alemu et al., 2011). Models estimating enteric CH$_4$ emissions can be broadly characterized as being empirical or mechanistic. Empirical models are based on mathematical or statistical associations of diet intake and composition and other animal factors with enteric CH$_4$ emissions. Mechanistic models are based on biochemical, metabolic, and physiological principles and attempt to simulate enteric CH$_4$ emissions on the basis of a mathematical description of fermentation biochemistry.
**Empirical Models.** Empirical models to predict CH₄ emissions have been developed since the 1930s (Kriss, 1931) and there are many models in this category found in the scientific literature. For example, Appuhamy et al. (2016) listed 40 such models that were developed in North America, Europe, Australia, and New Zealand. Because enteric CH₄ emissions are strongly related to feed intake, all models include a measure of intake, such as DMI, gross energy (GE) intake (GEI), ME intake, or NDFI. However, feed intake of individual animals is not routinely measured under commercial farm operations, and thus there may be a need to develop equations that do not require feed intake measures or estimates. The advantage of empirical models is that they can be constructed relatively easily from observed data and do not require a large number of inputs from the user. The most commonly used inputs for empirical model development are summarized in Figure 3. However, because enteric CH₄ emissions are affected by several factors other than feed intake, prediction ability may be compromised if the sample is not large enough and a representative population is not sampled. It is a challenge to represent CH₄-mitigating additives, including nitrate (Olijhoek et al., 2016) and 3-nitrooxypropanol (Hristov et al., 2015b) in existing empirical models. Empirical models are currently used to estimate the contribution of the livestock industry to GHG emissions, particularly enteric CH₄ emissions nationally and globally. For example, several countries, including the United States, use the following IPCC Tier 2 equation to determine enteric CH₄ emissions:

\[
\text{CH}_4 = Y_m \times \text{GEI},
\]

where CH₄ is enteric CH₄ emission in MJ/head per day, and Y_m = CH₄ conversion factor defined as percentage of GEI (MJ/head per day). This needs 2 kinds of inputs: feed DMI and the GE concentration of feeds. Although GE can be determined by bomb calorimetry, this analytical method is tedious and requires some expertise. Most forages and grains have a GE of approximately 18.4 MJ/kg of DM, but protein-rich or high-fat feeds such as oilseeds have a much greater GE as fats contain approximately 37 MJ/kg of DM and protein contains approximately 24 MJ/kg of DM, whereas feeds rich in minerals (ash) have a lower GE content. However, IPCC (1997) guidelines estimate GEI through determination of net energy requirements for body functions, which are then connected to DMI using estimated energy digestibility and digestible energy utilization efficiency. The steps involved in determining GEI and Y_m introduce errors in estimating enteric CH₄ emissions. The use of a constant value for Y_m is a major concern because it can vary considerably with varying DMI and DM digestibility (Appuhamy et al., 2016). It can take values ranging from 3 to 10% (Mills et al., 2003), and the IPCC Y_m constants do not encompass this range. Factors such as feed quality, production level (related to DMI), and diet composition affect the proportion of energy lost in the form of CH₄ (e.g., Moraes et al., 2014; Jayasundara et al., 2016). Hence, assigning a constant Y_m can lead to considerable uncertainty in the emission estimates, particularly in regions with diverse production systems. Several authors have challenged the use of constant Y_m value of 6.5 ± 1.0% of GEI (IPCC, 2006) across different regions of the world for dairy cattle (e.g., Kebreab et al., 2008). For example, the average Y_m for dairy cattle has been reported to be 5.4 to 5.7% for North America (Kebreab et al., 2008; Appuhamy et al., 2016; Jayasundara et al., 2016; Niu et al., 2018). The uncertainty around Y_m is about 1 percentage point, which is quite large and leads to gross overestimation of enteric CH₄ emissions for North America. In Europe, the Y_m varies between 6.0 and 6.9% (7.1 for Switzerland; Zeitz et al., 2012; Niu et al., 2018), and in Australia and New Zealand, the value is closer to the most recent IPCC recommendations at 6.6% (Appuhamy et al., 2016). Hence, recommendations for estimating enteric CH₄ emissions from dairy cows should be made on a regional rather than global basis. The analysis of Appuhamy et al. (2016) showed that no single empirical model is superior to others in all regions of the world. Any particular model may have strengths in simulating some aspects of the CH₄ emissions but not all at the same time. Multi-model ensemble methodology has become a widely accepted approach to improve prediction by taking advantage of complementary individual models and adjusting various biases, particularly in hydrology, climate, economy, and recently in crop growth models (Huang et al., 2017). If a regional or even global estimate of Y_m is desired, it may be possible to use the top 5 to 10 models within a region in a multiple CH₄ model ensemble to improve region-wide prediction.

**Mechanistic Models.** A limited number of mechanistic models have been developed to predict nutrient absorption from the digestive tract, including VFA, and these models have been modified to predict enteric CH₄ emissions by adding hydrogen calculations. These include the “Molly” model that describes nutrient utilization in cattle with the ability to predict enteric CH₄ emissions through hydrogen balance in the rumen (Baldwin, 1995); the “Cowpoll” model, which is based on a series of dynamic, deterministic, and nonlinear differential equations of nutrient utilization and includes CH₄ production in the rumen and hindgut (Dijkstra et al., 1992; Mills et al., 2001; Bannink et al., 2011);
the Nordic cow model “Karoline,” which is a dynamic, mechanistic model describing digestion and metabolism in dairy cows (Danfær et al., 2006; Huhtanen et al., 2015b); and the “AusBeef” model, which is a dynamic, mechanistic, and deterministic model of beef cattle production that uses a detailed representation of biological processes to determine nutrient utilization and CH₄ emissions (based on Nagorcka et al., 2000, which is an adapted version of the model of Dijkstra, 1994).

In all extant mechanistic models, the underlying principles in predicting CH₄ emissions are similar. The models predict nutrient digestion, absorption, microbial growth, and fermentation stoichiometry to determine type and amount of VFA production, hydrogen, and ultimately enteric CH₄ emissions during ruminal (and sometimes hindgut) fermentation. The models differ mainly in the number of microbial groups included, source and particle size of feed, substrates for VFA production, and VFA stoichiometry. Methane emissions are calculated in a similar way in all models, by calculating hydrogen balance in the rumen and assuming that any excess hydrogen is converted to CH₄. However, hydrogen production by cattle can be substantial, depending on diet composition, and hydrogen production shows large diurnal variation with peaks of production shortly after a meal (e.g., Hristov et al., 2015b; Guyader et al., 2015; Olijhoek et al., 2016; van Gastelen et al., 2017). Prediction accuracy of CH₄ emissions in mechanistic models depends largely on the accuracy of the stoichiometric models used and their accuracy to predict VFA molar proportions (Bannink et al., 2011). Alemu et al. (2011) evaluated several stoichiometric models and reported that their performance varies widely ranging from 5.2 to 43.2% RMSPE. There is a scarcity of studies that measured VFA production rates, because this requires the use of isotopes to differentiate between VFA concentrations (which are net production) observed in the rumen and production rates.

Researchers in the Netherlands apply a Tier 3 approach for national inventory of dairy cattle CH₄ emissions (based on country-specific experimental data and typically involving modeling and higher resolution land-use and land-use change data) using a mechanistic model (Bannink et al., 2011). Using this approach, Bannink et al. (2016) were able to explain part of the observed variation in enteric CH₄ emissions due to variation in grass silage quality, and DMI. Several model comparisons have been performed by Benchaar et al. (1998) and Kebreab et al. (2008), showing that the Cowpoll model agreed with observed data better than Molly or other empirical models for CH₄ emission from dairy cattle. For feedlot cattle, the Molly model performed better than Cowpoll [before Ellis et al. (2014) improved the prediction of CH₄ emissions and representation of rumen fermentation for finishing beef cattle]. Kass et al. (2017) compared the Molly model with Karoline model in their ability to predict CH₄ emissions and concluded that, although both models predicted CH₄ emissions reasonably well, the Karoline model was more accurate based on smaller mean and slope bias. The limitation

![Figure 3](https://example.com/figure3.png)

*Figure 3. Diet and animal factors used to estimate enteric methane production in extant empirical models. Color version available online.*
to the extensive use of mechanistic models of nutrient utilization is that they require inputs that may not be available at the production system level.

**Critical Data Gaps Limiting Enteric CH₄ Quantification**

Ellis et al. (2010) evaluated the prediction ability of several models to estimate enteric CH₄ emissions observed under various experimental conditions and concluded that, in general, predictions of these broadly applicable models were poor (based on RMSPE). According to Moraes et al. (2014), the poor predictive ability of current models can be due in part to the relatively small data sets used for model parameterization and the modeling techniques. Except for those that were developed by Moraes et al. (2014) for cattle in North America, most prediction models used a few hundred observations to develop relationships between enteric CH₄ emissions and dietary or animal factors. Normally, this number would not encompass the diversity of diets and animal factors in various regions of the world. Therefore, empirical models should ideally be developed from a database containing well over 1,000 individual observations or treatment means with accompanying information about dietary and animal factors that are known to affect enteric CH₄ emissions. In some cases, the improvement could be limited to the average animal, and observed variation might not be explained if key parameters are not included. Such databases will allow the development of robust estimates of average CH₄ emissions that can be tailored to be specific to a region and allow for various types of models ranging from simple one-covariate models to much more complex models that include several dietary and animal variables. Most of the CH₄ emission data in the literature originate from Europe, North America, Australia, and New Zealand (Appuhamy et al., 2016). Recently, there has been an increase in data being published from Central and South America (e.g., Dini et al., 2012; Muñoz et al., 2015), but there is still a dearth of data from Asia and Africa. Further research is required to produce data from indigenous and improved breeds of dairy cattle in Asia and Africa, and the data should encompass the production systems and feeds available in those regions. In our opinion, further development of enteric CH₄ prediction models would need regional data sets with as many data points as possible that have reliable DMI and CH₄ emission measurements. For the purpose of national inventories, the IPCC Tier 2 model with region-specific Yₜₐ factors would be most suitable.

Statistical methods that have been used in developing empirical models to date may not be appropriate because of the limitation of the framework used, such as not including random effects of animals or studies. Most of the current models (e.g., Ramin and Huhtanen, 2013) were developed with parametric inference gained from the likelihood function (frequentist statistical method). In this method, only a sequential application of simple significance tests can be calculated. In addition, only nested models can be compared, and different models are selected if alternative procedures or starting covariates are included in the statistical procedures. On the other hand, Bayesian methods are subjective and use prior beliefs to define a prior probability distribution on the possible values of the unknown parameters. Some examples of implementation of Bayesian modeling in animal nutrition and CH₄ emission prediction include those by Strathe et al. (2012) and Moraes et al. (2014). Even mechanistic, dynamic ruminant nutrition models used for enteric CH₄ emissions prediction can benefit from Bayesian methods to capture the inherent variability of the biological system under study and provide an assessment of the error associated with complex model results (Reed et al., 2016). Model evaluation methods have also advanced and it is possible to run Monte Carlo simulations and cross-validation techniques for large data sets and compare the predictive abilities of multiple CH₄ prediction models. Models should be developed at different complexity levels, which require different levels of activity data and dietary information for better functionality, as users will have various levels of information available to them in making predictions. In addition, the trade-off between model complexity and predictive ability should be quantified so users can decide whether the extra resources required for better prediction are justified by the increase in prediction. The trade-off has to be determined in the context of the aim for which the models are going to be used, such as for national inventory, assessment of mitigation options, and others.

Therefore, it is important that future models for a broad application be developed from large data sets with collaboration of scientists worldwide, as in the GLOBAL NETWORK project, and using robust state-of-the-art statistical techniques for model development and evaluation. The data sets should encompass a wide range of diets and production systems within regions and globally. It is also possible to develop a multi-model ensemble to improve enteric CH₄ emission prediction and determine uncertainty associated with the prediction.

**CONCLUSIONS**

There are large uncertainties in livestock CH₄ national and global inventories; sources of uncertainties in enteric CH₄ emission include animal inventories,
feed DMI, ingredient and chemical composition of the diet, and CH₄ emission factors. There is also significant uncertainty associated with enteric CH₄ measurements. Widely used measurement techniques are respiration chambers, the SF₆ tracer technique, and the GreenFeed system. All 3 methods need to be correctly and appropriately used to generate reliable and accurate data and valid tests of effects of diets and other treatments on enteric CH₄ emission or animal variation in CH₄ emission rates; some uncertainty remains as direct comparisons of techniques have shown inconsistent results. We emphasize that each of these techniques can have low accuracy and precision or produce misleading results if not properly implemented. Detailed guidelines for these techniques have been published and should be followed rigorously by researchers. Enteric CH₄ prediction models are based on various animal or feed characteristic inputs but are dominated by DMI in one form or another. Therefore, accurate prediction of DMI is of pivotal importance for accurate prediction of livestock CH₄ emissions. It is recommended that simplified enteric CH₄ prediction models based on DMI alone or DMI and limited feed- or animal-related inputs be developed and used for inventory purposes, where sufficient details or accuracy on dietary inputs are lacking. Broadly applicable and robust prediction models must be developed from large data sets generated through collaboration of scientists worldwide. To achieve high prediction accuracy, these data sets should encompass a wide range of diets and production systems within regions and globally. The uncertainty in enteric CH₄ prediction can be reduced by developing region-specific Y_m values. Similarly, the uncertainty in DMI estimation can be decreased by using DMI prediction equations that are region-specific instead of the GEI approach of IPCC Tier 2.

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