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Methane emissions from cattle: estimates from short-term measurements using a GreenFeed system compared with measurements obtained using respiration chambers or sulphur hexafluoride tracer

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Abstract

The GreenFeed (GF) system (C-Lock Inc., Rapid City, USA) is used to estimate total daily methane emissions of individual cattle using short-term measurements obtained over several days. Our objective was to compare measurements of methane emission by growing cattle obtained using the GF system with measurements using respiration chambers (RC) or sulphur hexafluoride tracer (SF_6). It was hypothesised that estimates of methane emission for individual animals and treatments would be similar for GF compared to RC or SF_6 techniques. In experiment 1, maize or grass silage-based diets were fed to four growing Holstein heifers, whilst for experiment 2, four different heifers were fed four haylage treatments. Both experiments were a 4 x 4 Latin square design with 33 d periods. GreenFeed measurements of methane emission were obtained over 7 d (days 22-28) and compared to subsequent RC measurements over 4 d (days 29-33). For experiment 3, 12 growing heifers rotationally grazed three swards for 26 d, with simultaneous GF and SF_6 measurements over two 4 d measurement periods (days 15-19 and days 22-26). Overall methane emissions (g/d and g/kg dry matter intake [DMI]) measured using GF in experiments 1 (198 and 26.6, respectively) and 2 (208 and 27.8, respectively) were similar to averages obtained using RC (218 and 28.3, respectively for experiment 1; and 209 and 27.7, respectively, for experiment 2); but there was poor concordance between the two methods (0.1043 for experiments 1 and 2 combined). Overall, methane emissions measured using SF_6 were higher ($P < 0.001$) than GF during grazing (186 vs. 164 g/d), but there was significant ($P < 0.01$) concordance between the two methods (0.6017). There were fewer methane measurements by GF under grazing conditions in experiment 3 (1.60/d) compared to indoor measurements in experiments 1 (2.11/d) and 2 (2.34/d). Significant treatment effects on methane emission measured using RC and SF_6 were not evident for GF measurements, and the ranking for treatments and individual animals differed using the GF system. We conclude that under our conditions of

use the GF system was unable to detect significant treatment and individual animal differences in methane emissions that were identified using both RC and SF₆ techniques, in part due to limited numbers and timing of measurements obtained. Our data suggest that successful use of the GF system is reliant on the number and timing of measurements obtained relative to diurnal patterns of methane emission.

Keywords: Dairy cattle, methane, respiration chamber, SF₆, GreenFeed

Abbreviations: CI, confidence interval; DM(I), dry matter (intake); GF, GreenFeed; LW, live weight; LWG, LW gain; NDIR, non-dispersive infrared; RC, respiration chambers; RFID, radio frequency identification; SF₆, sulphur hexafluoride tracer

1. Introduction

Accurate and robust measurements of methane emissions from individual animals are required for national inventories and assessment of mitigation strategies. There are a number of methods for determining methane emissions from ruminants, including respiration chambers (RC) and sulphur hexafluoride tracer (SF_6) techniques. Precise measurements of methane emission can be obtained by housing animals in RC, which allow direct measurement of total methane emission. However, RC are relatively expensive, have a limited throughput, and are disruptive to normal behaviour as animal by environment interactions that occur within grassland ecosystems are prevented. Respiration chambers are impractical for simulating grazing applications, and if the diet offered in the RC is fresh forage, then diet selection is limited, and eating patterns are likely to be determined by the feeding regime.

The SF_6 technique (Zimmerman, 1993; Johnson et al., 1994) can be used to make estimations of eructated and expired methane emissions from animals which can select their diet in a manner representative of farmed livestock (*e.g.* grazing). However, evaluations have challenged the precision of the SF_6 technique for estimating methane emissions (Vlaming et al., 2007; Pinares-Patiño and Clark, 2008; Pinares-Patiño et al., 2011), with greater between-animal variation compared to RC (Hammond et al., 2009). The SF_6 technique has also provided variable estimates of methane emission from animals on different herbages that have not been corroborated by RC measurements (*e.g.* Hammond et al., 2011; Waghorn et al., 2002; Sun et al., 2011 and 2012). Halter and collection canisters placed on the animal for methane estimates can interfere with grazing (Pinares-Patiño et al., 2008), especially with young animals, and a lower than predicted feed dry matter intake (DMI) will overestimate methane yields (g/kg DMI). Rumen SF_6 boluses must also be administered, and frequent

animal handling is needed, all of which can be disruptive to normal behaviour, and is relatively labour intensive.

In 2010, the commercial GreenFeed (GF) system (C-Lock Inc., Rapid City, South Dakota, USA) was introduced as a static short-term measurement device that measures methane emission from individual cattle, and uses head position sensors in combination with decision rules to assess the validity of measures obtained. The animal is free to move about and voluntarily enters a hood where a feed supplement (*i.e.* a reward for visiting the GF unit) is delivered. Measurements of methane emission by GF are typically over short (3-7 min) periods, several times within a day, over several days. The system is programmed using C-Lock Inc. software to control timing of feed availability and thus, encourage animals to distribute their voluntary GF visitation and hence methane measurements over a 24 h period so that ultimately a 24 h individual methane emission profile can be extrapolated from several days of short-term measurements. Cattle are typically not handled during GF operation and one GF unit can be used for numerous animals, with manufacturer recommendations of 15-20 animals/unit when grazing and 20-25 animals/unit if housed in free stalls. Because the GF system is relatively new, little is known about its operation, precision, accuracy, and the extent to which animal interaction with GF affects methane measurements.

The objectives of the present study were addressed across three experiments that included measurements of methane emission from individual growing dairy cattle using a single GF unit. Our objectives were to compare measurements of methane emission by growing dairy cattle obtained using GF with measurements using RC (experiments 1 and 2) and SF₆ (experiment 3). It was hypothesised that estimates of methane emission for individual animals and treatments would be similar for GF and RC or SF₆ techniques.

2. Materials and methods

Three experiments were used in this study whereby growing dairy cattle were fed a variety of diets and methane emission was measured using GF, RC and/or SF₆ techniques. Measurements in all experiments were individual DMI and methane production (g/d), calculated methane yield (g/kg DMI), and frequency of GF visitation (*i.e.* methane measurement frequency). All procedures used were approved and monitored under the UK Home Office Animals (Scientific Procedures) Act 1986.

2.1 Experiments

2.1.1 Experiment 1

Four Holstein Friesian dairy heifers aged 14 months with an initial live weight (LW) of 317 ± 20 kg were fed once daily (10:00 h) either maize or grass silage diets supplemented with or without an extruded linseed product (Lintec; 26% fat) at 6% of ration DM ($n = 4$ animals/treatment).

Experiment 1 was a 4 x 4 Latin square design with each period 33 d in duration, commencing with 21 d adaptation where access to GF was allowed, and GF data used for analysis was obtained during 7 d (days 22-28), after which animals were confined to RC for measurement of methane emission over 4 d (days 29-33).

Feed was offered to achieve target daily LW gains (LWG) of 0.75 kg. Feed intakes were measured on a daily basis using an electronic Calan Broadbent individual feeding system (American Calan Inc., Northwood, New Hampshire, USA) with feed refusals collected once daily before morning feeding. Animals were loose-housed and bedded on wood shavings with rubber mats and had *ad libitum* access to water.

2.1.2 Experiment 2

Four Holstein Friesian dairy heifers, aged 14 months with an initial LW of 339 ± 16 kg were fed twice daily (10:00 and 16:00 h in equal amounts), one of four conserved forage

(haylage) treatments of ryegrass, clover, trefoil and flowers ($n = 4$ animals/treatment). Further details of these treatments are given in Hammond et al. (2014). Similar to experiment 1, experiment 2 was a 4 x 4 Latin Square design with 33 d periods, with animals fed and housed in a similar manner as detailed for experiment 1.

2.1.3 Experiment 3

Twelve Holstein Friesian dairy heifers aged eight months, with a starting LW of 230 ± 6 kg, grazed the same treatments used to make haylage in experiment 2 (ryegrass, clover and flowers; $n = 12$ animals/treatment; Hammond et al., 2014). Heifers rotationally strip grazed each sward treatment for 26 d in a sequence of flowers, clover, then ryegrass. Each 26 d period commenced with 14 d adaptation where GF access was allowed, with simultaneous GF and SF₆ data obtained over two 4-d measurement periods (days 15-19 and days 22-26).

Dry matter intake was estimated using a rising plate meter (Farmworks Precision Farming Systems, Feilding, New Zealand) by taking 20 sward height readings before and after each days grazing period. Sward DM yield estimations were calibrated every second day by taking 5 x 0.5 m² quadrat cuts of the sward at a target post-grazing height of 6 cm and oven drying (100°C) the sample to give sward DM yield per m² which was applied to each sward height measurement.

2.2 Methane measurements

2.2.1 GreenFeed

The GF system measured methane emission using sensors that identified the animal and its head position within a sampling hood, air flow, and methane and carbon dioxide concentrations in exhaust air. GreenFeed operation was initiated when the animal placed its head inside the hood. A radio frequency identification (RFID) reader identified the animal's ear tag and GF sampling was activated when the animals head (located by an infrared sensor)

was in the correct location within the hood, and it was deemed that sufficient time had elapsed since the previous methane measurement for that animal.

Animal head position was critical for successful measurements as the animal is free to move its head in and out of the hood and thus only data captured with uninterrupted measurements was retained for statistical analysis. Position of the animals head within the hood was monitored using sensors to ensure complete breath collection. Adequate animal head position resulted in the dispensing of feed pellets which were used for enticement and encouraged the animal to maintain a suitable head position for accurate measurements. Pellets were dispensed from a hopper above the GF using a computer controlled rotating cup dispenser.

Animals were able to use the GF unit at any time, provided it was not in use by another animal, however, this did not necessarily generate a measurement of methane. A 'visit' is defined as a visit that results in a methane measurement. Thus, a 'visit' is only considered when a certain time has elapsed between visits (as dictated by the user) and a food reward is dropped, generating a methane measurement for that animal.

The concentration of the gas emitted by the animal was calculated using background gas concentration, the differential concentration of gas during the animal's time in the GF hood, and the calibration coefficient for concentration. The calibration coefficient was based on nitrogen, carbon dioxide and methane gases used to calculate the response of the sensors. The GF analysers were zeroed and calibrated weekly using zero baseline gas (oxygen-free nitrogen) and a span gas mixture nitrogen containing 5000 ppm carbon dioxide and 1000 ppm methane (BOC Ltd., Manchester, UK). This was to account for any drift in the calibration of the analysers, which was found to be negligible. A known amount of propane or carbon dioxide was released near where the animal's nose would be when feeding to check recovery

of expired gases when the physical location of the GF unit changed. There was no recovery correction required in the current study.

To measure gas production (mass per unit of time) an extractor fan was used to draw air past the animal's head into an exhaust pipe and airflow rate was measured. Airflow rate was multiplied by the increase in gas concentration when the animals head was in the hood. The duration the animals head was in the GF hood was recorded thus giving the time interval for calculation of mass per unit of time. The concentrations of methane and carbon dioxide gases were measured by non-dispersive infrared (NDIR) sensors, and an air filter was used to filter and remove any fine particulate material from the air that was subsampled to the sensors to prevent damage. The air filter was changed every two weeks. Data from GF was available real-time using mobile phone communication through a web-based data management system provided by C-Lock Inc.

For all experiments, the GF was programmed using C-Lock Inc. software to deliver a maximum of five rotations of a feed dispensing cup, delivering approximately 55 g of pellet (as fed) per rotation, with intervals of 45 sec between each rotation, so that 275 g of pellet was delivered during each visit. A maximum of four visits per day (24 h) was allowed, with a minimum of 4 h required between visits. Therefore, if an animal attempted to use the GF less than 4 h from the previous visit pellets would not be dispensed. Commercial calf pellets (Rearer18 Nuts, Wynnstay Group PLC, UK) were used for GF enticement and had a chemical composition (g/kg DM) of ash, 85.1; oil, 46.5; acid detergent fibre, 174; neutral detergent fibre, 289; starch, 259; water soluble carbohydrate, 91.3; nitrogen, 27.3, crude protein, 171; and gross energy (MJ/kg), 18.1. In all experiments total daily feed allocations included 1 kg of expected pellet DM provided by the GF unit.

The GF unit was set up indoors for experiments 1 and 2 at one end of animal housing, with gates positioned to restrict access to one animal at a time. Barn ventilation was used to

maintain ambient concentrations of methane in background air. For experiment 3, the GF unit was located outdoors under an awning at a point central to the experimental paddocks. The GF was located next to the only available water trough to encourage visitation, and fences and tracks were established to provide continuous access from grazed paddocks.

2.2.2 *Respiration chambers*

Details of the RC and measurements of methane emission are given by Reynolds et al. (2001) and Cammell et al. (1986). For measurements of gaseous emissions, two open-circuit RC were used (internal volume approximately 21 m³), with air-locks enabling access for faecal and urine collection (Cammell et al., 1986). An integrative sample of ambient and RC exhaust air was analysed at 4-min intervals, and every 4 h there was a switch to calibration gases (oxygen-free nitrogen and nitrogen carrier with 20.5%, 3000 ppm, and 200 ppm oxygen, carbon dioxide, and methane, respectively) to provide gas analyses with variation coefficients of 5% or less.

2.2.3 *Sulphur hexafluoride*

Experiment 3 used the SF₆ technique, as detailed previously by Hammond et al. (2014). Two weeks prior to experiment 3 commencing, heifers were each dosed by mouth with a SF₆ permeation tube (supplied by AFBI, Hillsborough, Northern Ireland, UK) into the rumen. The SF₆ gas release rates from the permeation tubes (5.176 ± 0.248 mg/d) were measured prior to dosing by oven incubation at 39°C and weighing twice weekly for six weeks. Daily methane emissions from heifers were estimated from analysis of air collected from around the nose and mouth over a 24 h period into a pre-evacuated PVC canister which was suspended under the neck. Based on recommendations given by Berndt et al. (2014), air was sampled using a crimped stainless steel capillary 0.004" ID 10 cm tube, with a flow rate between 0.45 to 0.55 ml/min. Canisters had a volume of approximately 2.3 L and a pre-collection vacuum of 90 kPa. Canisters were changed once daily at the same time each

morning and were rejected if vacuum post-collection was > 75 or < 50 kPa. A background air sample was also obtained daily from the paddock adjacent to that being grazed. Samples from canisters were analysed daily in our laboratory using gas chromatography to determine methane and SF₆ concentrations as described by Muñoz et al. (2012).

2.3 Data and statistical analyses

Data from GF and RC during periods 1 and 2 of experiment 1 were excluded from the analyses because the methane concentration of the calibration gas used for the GF unit was too low. Thus, comparisons for experiment 1 were restricted to periods 3 and 4 ($n = 8$) which meant treatment effects were not tested due to the limited observations obtained with the Latin Square design experiment. Each animal and period emissions data generated by GF was averaged over 7 d, whereas RC data was averaged over 4 d, with data expressed on a per min basis over 24 h and as a daily average (g/h and g/d).

For experiment 2, data from all four animals and periods were analysed statistically ($n = 16$) using the Mixed Models Procedures of SAS (2011) for random effects of animal and fixed effects of period and treatment. Like experiment 1, each animal and period emissions data generated by GF was averaged over 7 d, whereas RC data was averaged over 4 d in experiment 2, with data expressed on a per min basis over 24 h and as a daily average (g/h and g/d).

Experiment 3 provided methane data for 12 heifers grazing three fresh forage treatments for two 4-d methane measurement periods and two treatment periods (May to July and August to October; Hammond et al., 2014). Analysis of methane emission data were undertaken on daily averages across 4 d of measurements for both GF and SF₆ techniques (obtained simultaneously) for individual animals during each measurement period. Twelve heifers were used for the first grazing rotation of ryegrass, clover, and flowers, and also for

the second rotation of flowers. However, for the second rotation on ryegrass and clover, two animals were removed because there was insufficient sward cover. Therefore, a total of 136 GF and SF₆ individual animal average emission rate observations were analysed using Mixed Models Procedure of SAS (2011) for effects of forage treatment and treatment period (1 or 2), with 4-d measurement period within forage treatment period as a repeated effect within heifers (Hammond et al., 2014). When significant effects occurred, means of forage mixtures (clover and flowers) were differentiated from ryegrass control using Dunnett's adjusted mean comparisons.

Differences in methane emission (g/h, g/d and g/kg DMI) between techniques (GF vs. RC and GF vs. SF₆) across all experiments were tested using Lin's Concordance Correlation Coefficient analysis (Lin et al., 1998) in GenStat (2010) and the Univariate Procedure of SAS (2011) to determine if the difference between the two methods for each experiment was different from zero.

Within each experiment, the Least Squares Mean option of the GLM procedure (SAS, 2011) was used to rank individual animals according to their methane emission (g/d and g/kg DMI) for each measurement technique using animal as a fixed effect. In addition, the GLM procedure was used to regress GF measurements against RC or SF₆ measurements (g/d).

3. Results

3.1 General observations

3.1.1 Experiment 1

As stated previously, experiment 1 included data from four animals with $n = 8$ observations (only two periods out of a possible four were used). Dry matter intake during GF and RC measurements was similar (Table 1). Average methane production (g/d) and yield (g/kg DMI), determined using either GF or RC techniques, was similar for individual animals

(198 vs. 215 g/d, and 26.6 vs. 28.3 g/kg DMI, for GF vs. RC techniques, respectively) (Table 1). Individual animals had a similar methane output regardless of measurement technique used, however methane data (g/d and g/kg DMI) generated by GF and RC techniques ranked heifers differently in numerical order from high to low methane output (data not shown).

3.1.2 Experiment 2

There were four heifers used in experiment 2 with 16 observations (all four periods included). Heifers had a similar DMI during GF and RC measurements (Table 1). Average daily methane production (g/d) and yield (g/kg DMI) did not differ with measurement technique for individual animals (208 vs. 209 g/d, and 27.8 vs. 27.7 g/kg DMI, for GF vs. RC techniques, respectively; Table 1). For both GF and RC methods, animals were significantly ($P = 0.05$) different to each other in their methane production but not methane yield. Both GF and RC techniques ranked animals in numerical order, from low to high, the same for methane production, but not for methane yield (data not shown).

3.1.3 Experiment 3

Experiment 3 used 12 heifers and had 136 observations. Approximately 88% of SF₆ canisters were accepted (478 measurements out of a possible 544), with 12% of measurements unsuccessful due to sampling tube blockages, broken collection tubes, or displacement of canisters from the heifer. Both GF and SF₆ techniques were used simultaneously so DMI was the same with measurement technique. Daily methane production determined by GF for individual heifers was lower ($P < 0.001$) than SF₆ (164 vs. 186 g/d, respectively; Table 3). For both GF and SF₆ methods, heifers were significantly ($P = 0.05$) different to each other in their methane production (g/d), and the ranking of animals, from low to high methane production, was different for the two techniques.

Insert Table 1 here

3.2 Technique comparisons

3.2.1 GreenFeed vs. respiration chamber

Combining data from experiments 1 and 2, Lin's Concordance Correlation Coefficient between GF and RC, when used to measure methane production and yield of individual heifers, was 0.1043 and 0.058, respectively, with a non-significant ($P > 0.50$) association between the two techniques, based on the 95% confidence interval (CI) (Fig 1). There were diurnal patterns of methane erucation over a 24 h period for animals in both experiments 1 and 2, measured using GF and in RC (Fig 2). Emissions ranged from about 4 g/h immediately before their morning feeding to a maximum of about 15 g/h after feeding, on both silage and haylage diets. The increase in methane production after 10:00 h in experiment 1 relates to once daily feeding, whereas the increases just after 10:00 and 16:00 h represent the twice daily feeding regime. Based on all methane measurements, compared to the GF data, there was less variability with the RC emission measurements (g/d) from both experiments, in part because measurements for GF were much less frequent and fewer in number than for RC (Fig 2).

Insert Fig 1 here

Insert Fig 2 here

3.2.2. GreenFeed vs. sulphur hexafluoride

Lin's Concordance Correlation Coefficient between GF and SF₆ techniques, used to measure methane production from individual heifers of experiment 3, was 0.602, with a significant ($P < 0.01$) association between the two techniques, based on the 95% CI (Fig 3).

Insert Fig 3 here

3.3 GreenFeed for detecting dietary treatment effects

In experiment 2, DMI and methane production during RC measurements was affected by haylage type ($P = 0.045$ and $P = 0.025$, respectively), but this was not evident for GF measurements (Table 2). When methane was expressed in terms of DMI (yield, g/kg DMI), RC detected significant differences ($P = 0.020$) between haylages, but GF did not. There was no consistency in the relative difference between measurement techniques with dietary treatment. Relative to RC, GF underestimated 15% of methane yield when heifers were fed a ryegrass diet, compared to an overestimation of 12% for heifers on a flower diet (Table 2).

For heifers of experiment 3, methane production (g/d) differed significantly with both GF and SF₆ techniques ($P = 0.019$ and $P < 0.001$, respectively) for all three forage treatments. However, the ranking of mean estimates for the different forages differed with technique (Table 2). When methane was expressed in terms of DMI (methane yield), noting that the techniques estimated methane simultaneously, the ranking of treatment means was not the same for GF ($P = 0.080$; flowers > clover = ryegrass) and SF₆ techniques ($P = 0.002$; clover = ryegrass > flowers). For two out of three dietary treatments fed, GF underestimated methane yield relative to SF₆ by up to 18% (Table 2).

Insert Table 2 here

3.4 GreenFeed visitation

During the 14 d of GF measurements in experiment 1, there were a total of 118 visits to the GF unit, averaging 2.11 visits/d. For the 28 d measurement period in experiment 2, total GF visitation was 262, averaging 2.34 visits/d. During the 48 d of measurements for experiment 3, heifers visited the GF unit 880 times, averaging 1.60 visits/d (Table 3). The

average duration (min:sec) of GF measurements for experiments 1, 2, and 3 were 04:44, 04:43, and 04:58, respectively.

Figure 4 shows the pattern of visits to the GF, according to hour of the day. For all experiments, animals frequented the GF most often between 07:00 and 08:00 h, and between 13:00 and 14:00 h, with fewer visits in early morning hours (between 01:00 and 06:00 h). GreenFeed measurements were prevented if another animal was already using the unit, when animals were yarded for other experimental activities such as SF₆ canister changes, and during the allocation of new grazing. The type of diet offered affected GF visitation for experiment 3, but not experiment 2 (Table 2). Heifers in experiment 3 made fewer ($P < 0.001$) visits overall to the GF when on the ryegrass (219) and clover (229) paddocks, compared to the flower (432) paddock.

Insert Table 3 here

Insert Fig 4 here

4. Discussion

4.1 Comparison of measurement techniques

4.1.1 GreenFeed vs. respiration chamber

Based on the concordance analysis for methane emission from heifers of experiments 1 and 2, GF and RC techniques had a poor agreement, yet average methane emission overall was similar for the two techniques. It is difficult to interpret these conflicting results; however the large amount of variation about the line of equality (Fig 1) is a likely explanation for overall methane means being similar between techniques (Table 1) but having low concordance correlation. The lack of concordance between methods is in part attributable to the relatively small number of short-term measurements obtained by GF on each day of

measurement. The concept behind the GF system is that although it is unknown what an animal is eructating when not visiting the GF, the accumulation of data over 24 h can provide a representative pattern (Fig 2). Thus, the GF technique relies on the animal visiting the unit at different times during the day to characterise the daily pattern of methane emission over a number of days. In contrast, RC measurements in this study were based on integrated measurements every 4 min over four consecutive days.

The inability of GF to detect changes in methane production due to treatment or animal effects compared to RC (and SF₆) is not unexpected given the methodology the technique employs. Enteric methane production from ruminants typically exhibits a diurnal pattern related to feeding and meal consumption, with methane emission rate varying by as much as five-fold over the course of a day (Crompton et al., 2010). Peak enteric methane production occurs approximately 120 and 60 min after the morning and afternoon feeds, respectively, for a lactating dairy cow fed *ad libitum* twice daily (Crompton et al., 2010). Frequent or continuous measurements over a 24 h period using RC or SF₆ account for any diurnal variation in methane production, but intermittent short-term measurements may vary significantly depending on when those measurements are taken during the day.

There was a greater range in absolute emissions for both measurement techniques with experiment 2 data compared to experiment 1 that was associated with greater differences in DMI and methane production. However, when emissions were expressed per unit of feed intake (g/kg DMI), relationships between GF and RC measurements were still weak and variation was large, especially for the GF measurements. The variable relationship suggests that the differences in methane emission due to treatments and animal variation measured by the RC are not correlated with differences measured by GF. In other words, ranking of the animals according to methane production and yield differed between the two techniques, despite substantial differences being observed. The absence of a significant correlation

between GF and RC measurements for individual animal observations (Fig 1) casts doubt on the capability of GF to distinguish (and rank) individual animals under the conditions with which GF was used in our experiments. With the exception of daily methane production in experiment 2, GF and RC ranked heifers differently in their methane emission. Daily mean methane production varied from about 160 to 270 g/d measured in RC, and although GF also recorded a similar range, the range was for different heifers on different diets (data not shown).

It is possible that the algorithms used by the GF system for the calculation of methane output, or the timing of visits relative to daily patterns of methane emission, may account for the discrepancies observed between GF and RC data. GreenFeed calculations are based on differences in the concentration of the air exhaled and eructed by the animal, less background air concentrations measured pre- and post-feeding. The GF is able to differentiate emissions of methane in exhaled air above background, so exhaled air is included in the emission calculation. The calculations are reliant on erucation events taking place within the measurement period, and the algorithms may need to be modified to increase accuracy and reduce variation. For the animals and diets used in our study, more GF measurements were needed over a longer period, and at more frequent intervals, to better represent the diurnal pattern of methane emissions over 24 h. Increased animal visitation to the GF may require longer periods of measurement (more days), more visits per day (and thus greater feed consumption), or the use of an alternative ‘bait’ (Hegarty, 2013). In addition to this, it has been estimated that 1-2% of methane is voided as flatus (Murray et al., 1976), and it will contribute to methane emissions (Ellis et al., 2008) measured in RC. These considerations for GF measurements are also pertinent to other on-farm breath analysis techniques (*e.g.* Garnsworthy et al., 2012).

4.1.2 *GreenFeed* vs. *sulphur hexafluoride*

GreenFeed and SF₆ techniques had moderate concordance (agreement), in part due to the greater number of observations compared. However, overall methane emissions determined using GF were significantly lower than those measured using the SF₆ technique. Differences between GF and SF₆ techniques are likely due to the duration of methane measurements obtained for each animal. The SF₆ technique is based on integrative sampling with a sampling duration of nearly 1440 min/d (100% of 24 h). In comparison, the GF unit is designed to take intermittent samples, and based on the average of the three experiments presented here, sampling duration (5 min/visit at 2 visits/animal/d) was only 10 min/d (0.7% of 24 h).

All tracers have weaknesses (Shipley and Clark, 1972) and the variation associated with SF₆ estimates may be in part a consequence of factors affecting the technique itself (Deighton et al., 2013, 2014a and 2014b), or alternatively the variation might be real. Recent work has found that successful use of the SF₆ technique to detect differences in enteric methane emissions due to diets or between animal species may be confounded by diet or genetic effects on body temperature (Deighton et al., 2014b). In order to accurately determine methane emissions, it is necessary that gases are collected continuously at a constant rate for 24 h, however; it has been recently shown by Deighton et al. (2014a) that capillary tubes are unsuitable for use as flow restrictors to achieve this, causing a bias of up to 15.6% in calculated methane emissions. Deighton et al. (2014a) has since proposed a ‘modified SF₆ technique’ which incorporates orifice plate flow restrictors for 24 h sample collection and has found technique error associated with SF₆ release, sample collection and analysis to be reduced.

4.2 *GreenFeed* for detecting dietary treatment effects

Although all three techniques measured significant treatment effects on methane emissions, the ranking of these effects differed with measurement technique. Critically, both the RC and SF₆ techniques found methane yield (g/kg DMI) to be the lowest in both experiments 2 and 3 for animals fed flowers compared to the other dietary treatments (Hammond et al., 2014). GreenFeed on the other hand, was unable to detect treatment effects on methane yield in experiment 2, and ranked the treatments differently to SF₆ for experiment 3. This in part reflects the variability of GF measurements attributable to the timing and limited number of short-term measurements obtained in the present experiments.

4.3 GreenFeed visitation

Although animals had few problems adapting to the GF and used it willingly, visits were less frequent than permitted, particularly for grazing animals in experiment 3. The lack of GF visits from animals both while out grazing and during early morning, may have negatively biased methane production measured by GF (Fig 3). The low frequency of visits between 09:00 and 13:00 h (Fig 4) is likely to be when peak methane emissions occur in a once daily feeding system, as can be seen from the rise in methane production (g/h) in Fig 4. Thus, the infrequent daily measurements made by the GF system in experiment 1 is a likely explanation for numerically lower methane emissions from the GF compared to RC. In experiment 2, the GF pattern of visitation was better distributed over the course of the day, although a weaker relationship occurred between the two techniques (Fig 2).

In all experiments, fewer visits occurred in the early morning hours, and a lack of methane data over this period may have affected the average estimate of daily emissions. Every morning, heifers of experiment 3 were given a new allocation of feed at about 10:30 h after SF₆ canisters were replaced. The allocation of new pasture is likely to have been responsible for the drop in GF visits between 09:00 and 13:00 h. During this period of time,

methane emissions would have been at their highest, which must partly account for the 13% greater daily methane emissions determined by the SF₆ technique, compared to the GF system. The lower visitation to the GF by grazing heifers is cause for concern, especially as the GF system relies on having enough daily measurements over the course of an ‘average’ day to estimate daily emissions. Further evaluations of the GF system should determine the number of days and measurements per day required for GF to provide accurate and precise measures of methane emissions.

It would appear that the number of visits to the GF is influenced by dietary treatment (and possibly level of feed intake), with more visits made when heifers were grazing flowers, compared to ryegrass and clover (experiment 3). For experiment 3, this may have been attributable to the location of the GF relative to the paddocks, as well as the DM and nutrient content of the swards grazed. It would appear that less favourable diets may contribute to increased GF visitation when a favourable ‘treatment’ *i.e.* pelleted concentrates, is rewarded, and this may have consequences for methane estimates on different treatments (different numbers of samples and sampling times for each treatment). This is a concern for nutrition experiments that investigate effects of diet composition on methane emission if diet comparisons are affected by varying amounts of feed reward provided by GF.

One concern with the use of the GF under our conditions is the temporal distribution of GF visitation and the potential for bias in methane emission measurements by the GF system. This is because unlike both RC and SF₆ techniques the methane measurement obtained from each individual animal by the GF system is voluntary and thus not completely independent. The use of the GF unit by each individual animal within the group, and therefore the temporal distribution of their methane measurement, is affected by their cohorts and environmental circumstances. The inclusion of a given animal in the GF unit causes the exclusion of all other animals within the group. Strictly speaking an individual animal is not

totally independent as an experimental unit when the GF system is used and therefore to achieve completely independent experimental replication for individual animals housed as groups in pens or paddocks multiple GF units may be required.

5. Conclusions

Overall, the GF system provided an estimate of methane emission by growing dairy cattle that was not different from RC measurements, but significantly lower than for SF₆. However, concordance analyses found no agreement between GF and RC, and only moderate agreement with SF₆. We conclude that as used in our experiments, the GF system was unable to detect significant treatment and individual animal differences in methane emissions that were identified using both RC and SF₆ techniques. The successful use of the GF system is reliant on the number and timing of measurements obtained relative to diurnal patterns of methane emission. Therefore, animal and diet type, intake level and appetite (*e.g. ad libitum* vs. restrictive feeding), total feed availability, accessibility of the GF unit relative to other feeds and activities, as well as type, amount, and timing of feed used to elicit GF use all affect GF visitation and thus measurements of methane emission using the GF system. Multiple animals using a GF unit can alter the temporal distribution of measurements for individual animals and this potential bias should also be considered in designing future experiments.

Further evaluation of GF is needed to determine how best to deploy the system to meet specific objectives, the number and timing of measurements required for specific measurement conditions, as well as the capacity of the GF to detect significant changes in methane emissions with individuals and treatments. We suggest that in addition to increased frequency of daily GF visits future studies should include longer periods of measurement and a greater number of animals per treatment than is required for RC studies.

534

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541

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Table 1 Dry matter intake (DMI; kg/d), methane production (g/d), and methane yield (g/kg DMI) from growing dairy cattle within three different experiments using GreenFeed (GF), respiration chamber (RC) and sulphur hexafluoride tracer (SF₆) techniques.

	Experiment 1 ^a	± SD	Experiment 2 ^a	± SD	Experiment 3 ^b	± SD
DMI, kg/d						
GF	7.62	0.81	7.60	0.81	9.15	2.67
RC ^a or SF ₆ ^b	7.66	0.59	7.54	0.94	9.15	2.67
<i>n</i>	8		16		136	
SEM	0.132		0.182		N/A	
P	0.799		0.747		N/A	
Methane, g/d						
GF	198	20.4	208	31.5	164	51.0
RC ^a or SF ₆ ^b	215	22.3	209	30.9	186	57.3
<i>n</i>	8		16		136	
SEM	9.230		10.59		2.900	
P	0.170		0.940		< 0.001	
Methane, g/kg DMI						
GF	26.6	2.80	27.8	5.62	18.8	6.94
RC ^a or SF ₆ ^b	28.3	3.01	27.7	1.81	21.5	7.60
<i>n</i>	8		16		136	
SEM	1.365		1.459		0.349	
P	0.255		0.933		< 0.001	

^a Experiments 1 and 2 used RC for measurement of methane from dairy heifers.

^b Experiment 3 used SF₆ for estimate of methane from grazing dairy heifers.

^c DMI was measured using Calan gates for individual animals in experiments 1 and 2, however for experiment 3, DMI was estimated by pre- and post-herbage mass (hence same DMI for animals where both GF and SF₆ were used simultaneously).

Table 2 The difference in methane emission between GreenFeed (GF), respiration chamber (RC), and sulphur hexafluoride tracer (SF₆) techniques with dairy heifers fed different dietary treatments.

Experiments	<i>n</i>	Dry matter intake (DMI), kg/d ^c		Methane production, g/d		Methane yield, g/kg DMI			Relative difference between methods (%)
		GF	RC ^a or SF ₆ ^b	GF	RC ^a or SF ₆ ^b	GF	RC ^a or SF ₆ ^b	Difference ^d	
Experiment 2^a									
Ryegrass	4	8.28	8.13	196	230	24.1	28.4	-4.32	-15
Clover	4	6.86 ^d	7.10 ^b	202	200 ^c	29.5	28.1	1.40	5
Trefoil	4	7.93	7.51	226	218	28.9	29.2	-0.32	-1
Flowers	4	7.34	7.42 ^c	209	190 ^b	28.8	25.7 ^c	3.14	12
SEM		0.377	0.255	17.33	8.890	3.013	0.662	2.844	
P (haylage)		0.180	0.045	0.515	0.025	0.521	0.020	0.298	
Experiment 3^b									
Ryegrass	44	10.0	10.0	175	204	17.3	21.8	-3.38	-16
Clover	44	8.69 ^a	8.69 ^a	166	202	18.5	23.0	-4.24	-18
Flowers	48	8.78 ^b	8.78 ^b	159 ^b	159 ^a	19.7 ^c	19.5 ^c	0.48	2
SEM		0.230	0.230	5.420	4.989	0.768	0.734	0.754	
P (forage)		0.001	0.001	0.019	< 0.001	0.080	0.002	< 0.001	

For each parameter, different letters indicate significant differences from the ryegrass control according to Dunnetts test (^a P < 0.001, ^b P < 0.01, ^c P < 0.05, ^d P < 0.10).

^a Experiment 2 used RC for measurement of methane from dairy heifers

^b Experiment 3 used SF₆ for estimation of methane from grazing dairy heifers

^c DMI was measured using calan gates for individual animals in experiment 2, however for experiment 3, DMI was estimated by pre- and post-herbage mass (hence same DMI for animals in experiment 3 with measurement technique)

^d Difference is generated using GF value less corresponding RC or SF₆ value

659 **Table 3** Animal visitation to the GreenFeed (GF) unit across three different experiments

660

	Number of measurement days	Total number of GF visits (methane measurements)	Total number of GF visit (methane measurements) per animal per day
Experiment 1			
Total	14	120	2.11
SD			0.49
SEM			0.17
Experiment 2			
Ryegrass		76	2.71
Clover		60	2.14
Trefoil		68	2.43
Flowers		58	2.07
Total	28	262	2.34
SD			1.05
SEM			0.26
P (haylage)			0.425
Experiment 3*			
Ryegrass		219	1.24
Clover		229	1.30 ^d
Flowers		432	2.26 ^a
Total	48	880	1.60
SD			1.09
SEM			0.07
P (forage)			< 0.001

* For experiment 3, different letters indicate significant differences from ryegrass control according to Dunnetts test (^aP < 0.001, ^dP < 0.10)

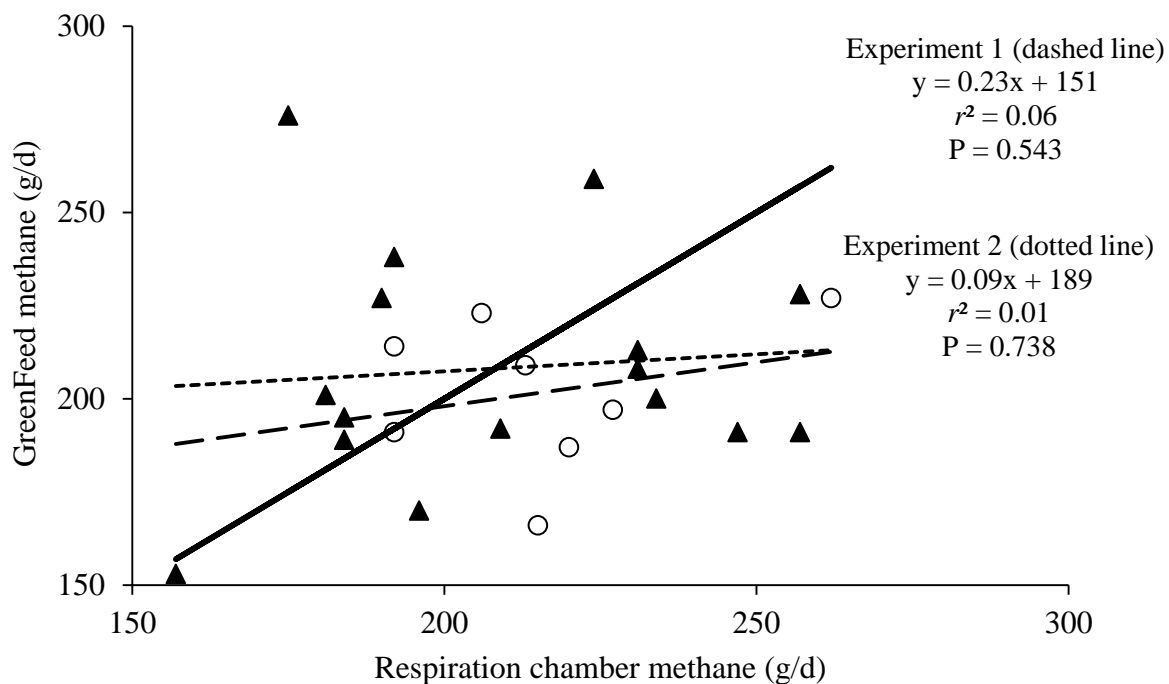


Fig 1. Relationships between methane production (g/d), determined using GreenFeed (GF) and respiration chamber (RC) techniques, of individual dairy heifers in experiments 1 (open circle symbol; $n = 8$) and 2 (closed triangle symbol; $n = 16$). Solid line indicates $y = x$. Lin's Concordance value for both experiments combined = 0.1043.

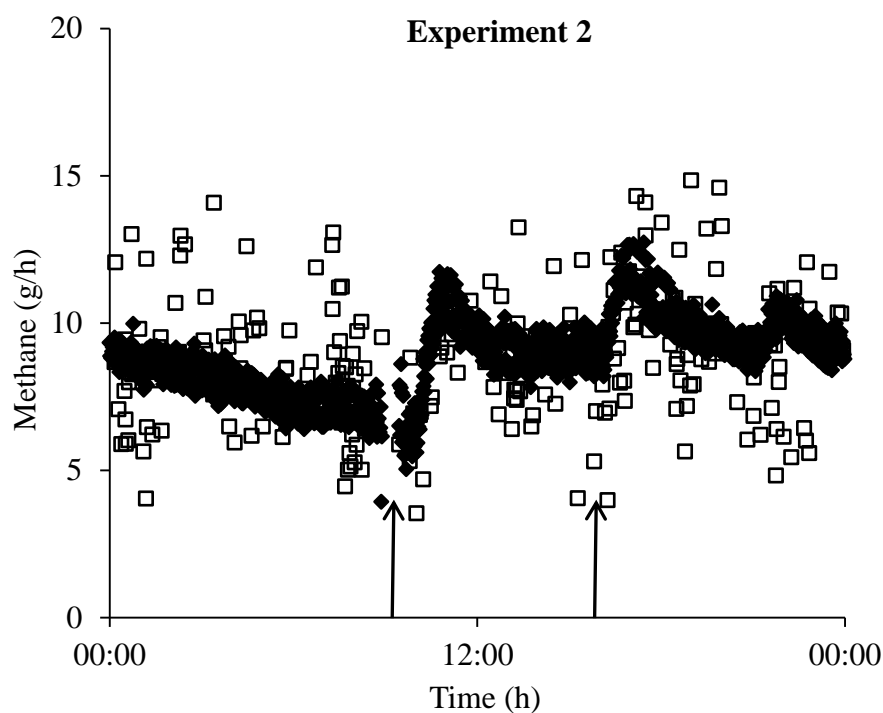
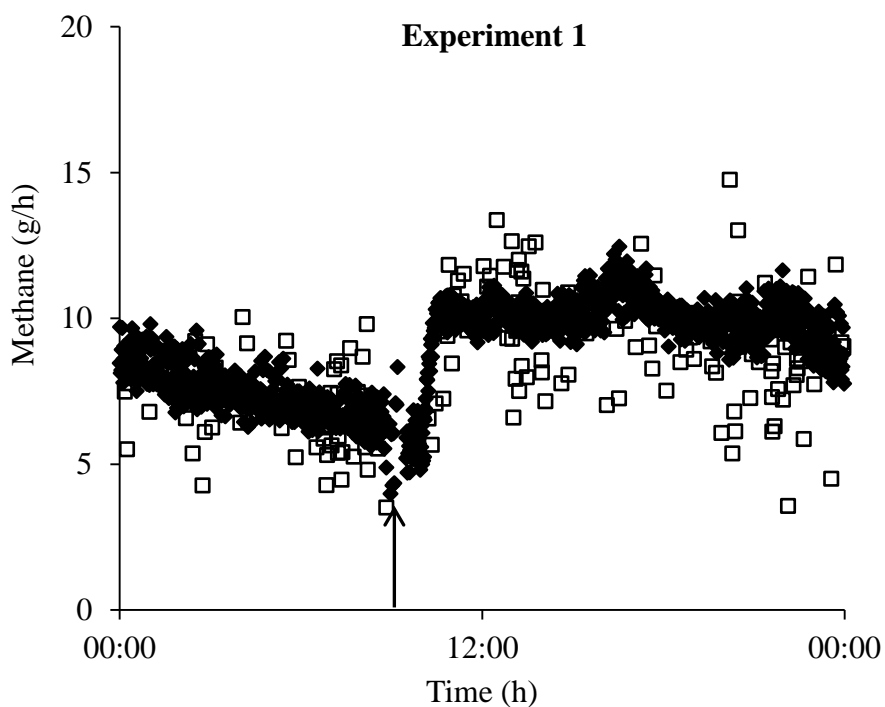
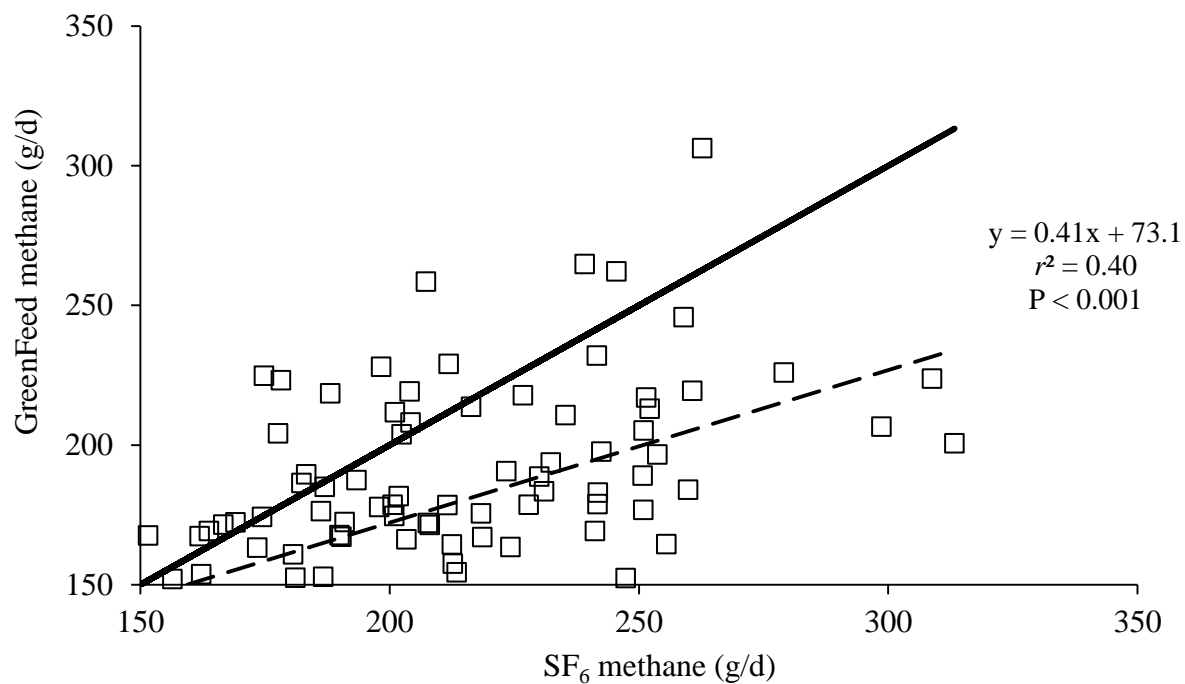
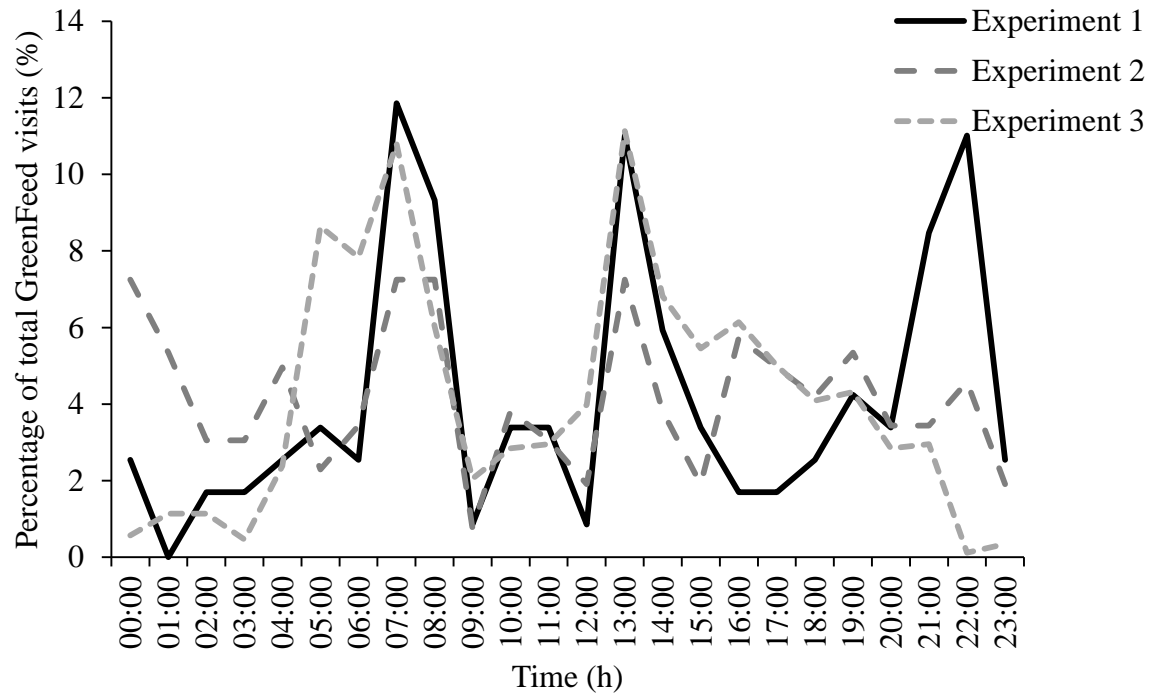


Fig 2. Comparison of methane emission rate (g/h; minute average) measured using GreenFeed (GF; open square symbol) and respiration chambers (RC; closed diamond symbol) for all dairy heifers in experiments 1 ($n = 8$) and 2 ($n = 16$). There were 56 d GF and 32 d RC measurements for experiment 1, and 112 d GF and 64 d RC for experiment 2. Arrows indicate time of feeding.



674

675 **Fig 3.** Relationship between methane production (g/d), determined using GreenFeed (GF)
 676 and sulphur hexafluoride tracer (SF₆) techniques, of individual dairy heifers in experiment 3
 677 ($n = 136$). Solid line indicates $y = x$. Lin's Concordance value = 0.6017.



678

679 **Fig 4.** Diurnal pattern of GreenFeed (GF) visitation (methane measurements) over 24 h, as a
 680 percentage of total visits, by growing dairy cattle of experiments 1 (120 GF visits/14 d), 2
 681 (262 GF visits/28 d) and 3 (880 GF visits/48 d).