Inclusion of a tannin-rich legume in the diet of beef steers reduces greenhouse gas emissions from their excreta

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The objectives of this study were to determine the emission of nitrous oxide (N₂O), methane (CH₄), and carbon dioxide (CO₂), as well as the isotopic composition of N₂O from excreta of beef steers fed ‘AU Grazer’ sericea lespedeza hay [SL; Lespedeza cuneata (Dum. Cours.) G. Don]. Fifteen Brahman × Angus crossbred steers were fed one of three experimental diets: 0, 50, or 100% inclusion of SL into ‘Tifton 85’ bermudagrass hay (Cynodon spp.). Gas sampling occurred on days 0, 1, 3, 5, 7, 14, 18, 25, and 32 after urine or feces application to static chambers for two experimental periods. Effect of the day after feces application (P < 0.001), while day × inclusion of SL interaction was observed in urine (P < 0.001) for all greenhouse gases (GHG) analyzed. Peaks of emission of all GHG in urine and feces occurred in the first days (P < 0.001), with days 3 and 5 being most depleted in 15N-N₂O in feces, and days 3, 5, and 7, in urine (P < 0.001). Feeding SL to beef steers was effective in mitigating the emission of GHG from the excreta, but further research is necessary to investigate the mechanisms behind the reductions.

Anthropogenic activity has modified natural environmental processes by increasing atmospheric concentrations of the main greenhouse gases (GHG), including through the expansion of agricultural activities1. In 2021, world cattle inventory was reported at 1 billion head2, with both enteric fermentation and manure being major contributors of methane (CH₄) and nitrous oxide (N₂O)3.

Manure is a significant source of GHG in grasslands due to the presence of organic compounds and their decomposition under anaerobic conditions4. Anaerobic bacteria decompose the organic material releasing CH₄, whereas interactions of different sources of N affect the N cycle, thereby, influencing the daily fluxes of N₂O5. The main concern is that the amount of N in the excreta deposited in a specific area exceeds the immediate plant needs, and the excess N is lost through nitrate leaching and N₂O6,7.

Nutritional adjustments are a strategy to modify the intensity and frequency of the processes that lead to the generation and emission of GHG in the excreta of ruminants. Animal diet, as well as the feed quality and quantity can influence urine and feces N concentrations. Forage legumes, for example, provide a wide range of secondary metabolites, such as tannins, that can modulate ruminal fermentation6. Condensed tannins (CT) bind to dietary protein protecting it from ruminal degradation, increasing the uptake of amino acids by the small intestine and the excretion of N in the feces7. Moreover, while natural nitrogenous compounds in the urine of cattle were reported to inhibit N₂O formation processes in the soil10–12, tannins would affect the proportion of those compounds, which can consequently affect the emissions of N₂O13.

Static chambers are a commonly used technique to measure GHG fluxes from soils, but due to the heterogeneous nature of the fluxes14, there are uncertainties in the magnitude, distribution, and temporal pattern of the natural and anthropogenic sources. This limits the accuracy of the results and comparisons among studies.

References

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from different locations. Within this context, the analyses of the natural abundance of stable isotopes from the gases in the atmosphere can represent a powerful ally to estimate GHG budgets, since the processes of GHG formation and sequestration have specific ratios of heavy to light isotopes.

We hypothesized that the inclusion of a tannin-rich legume in the diets of beef steers would decrease the emission of N\textsubscript{2}O, CH\textsubscript{4} and CO\textsubscript{2} from their excreta, due to the shift in N excretion from urine to feces and to the impact of tannins on methanogenesis. Additionally, urine and feces would present different emission factors and isotopic composition of N\textsubscript{2}O, enabling source attribution to the emissions. Therefore, the objectives of this study were to estimate the net emissions of N\textsubscript{2}O, CH\textsubscript{4}, and CO\textsubscript{2} from excreta of beef steers fed a tannin-rich forage legume, as well as to determine the emission factor of N\textsubscript{2}O and isotopic composition of N\textsubscript{2}O.

Results

N\textsubscript{2}O flux and δ\textsuperscript{15}N isotopic composition. There was an effect of sampling day on N\textsubscript{2}O emissions from feces, with the first 5 days presenting greater emissions compared to rest of the days (P < 0.001), and intermediate emissions at day 14 (Fig. 1a). For urine, an interaction between SL inclusion level and day of N\textsubscript{2}O emissions was observed (P < 0.001), with emissions being significantly greater on days 1, 3, and 5 after urine application (P < 0.001), and treatment 100SL presented the least emissions.

Peak N\textsubscript{2}O emissions in feces occurred 5 days after its application for all diets (0.12, 0.07, and 0.08 mg m\textsuperscript{-2} d\textsuperscript{-1} for 0SL, 50SL, and 100SL, respectively) and decreased to similar background emissions after 18 days (0.05 mg m\textsuperscript{-2} d\textsuperscript{-1}, P > 0.05). On the other hand, peak N\textsubscript{2}O emissions in urine occurred earlier (1 or 3 days after application; 0.08, 0.06, and 0.05 mg m\textsuperscript{-2} d\textsuperscript{-1} for 0SL, 50SL, and 100SL, respectively) but took longer (25 days) to reduce to background emissions (0.02 mg m\textsuperscript{-2} d\textsuperscript{-1}, P > 0.05), despite no significant differences among SL levels being observed from day 7 onward.

There were effects of day (P < 0.001) on urine and feces δ\textsuperscript{15}N composition from N\textsubscript{2}O (Fig. 2). The most δ\textsuperscript{15}N depleted N\textsubscript{2}O occurred on days 3 and 5 for both types of excreta (− 4.2 and − 8.2 ‰ for feces, and − 2.2 and − 5.2 ‰ for urine).

CH\textsubscript{4} and CO\textsubscript{2} emissions. There were effects of sampling day on CH\textsubscript{4} and CO\textsubscript{2} emissions from feces (Figs. 3a and 4a, respectively; P < 0.001) and SL inclusion × day interaction on emissions from urine (Figs. 3b and 4b, respectively; P < 0.001). For CH\textsubscript{4}, the peak in feces occurred on the first day after the application of excreta, with
no significant differences among levels of SL inclusion observed for the other days. In urine, CH₄ emission was different among the levels of SL only on the first day after its application, with greatest emission demonstrated by treatment 0SL (0.0006 mg m⁻² d⁻¹) and least, by treatment 100SL (−0.0004 mg m⁻² d⁻¹). Additionally, negative CH₄ peaks, regardless of the level of SL inclusion, were observed on day 18 after application of urine, which was restored to background levels on day 25.

Peak CO₂ in feces occurred on day 1 after the application of excreta (P < 0.001), regardless of the level of SL, and reductions to background levels were reached at day 18 (Fig. 4b). In urine, although peak was observed on the same day (day 1) for treatment 100SL, peak CO₂ for treatments 0SL and 50SL occurred 5 or 14 days after excreta application, respectively.

Excreta composition, cumulative emissions of GHG and N₂O emission factor. Nitrogen concentration observed in feces was 1.6, 2.2, and 2.8%, while in urine, it was 0.19, 0.24, and 0.58% for 0SL, 50SL and 100SL, respectively (Table 1).

Cumulative emissions of N₂O from feces were 2.02, 1.09, and 0.84 mg m⁻² for 0SL, 50SL, and 100SL, respectively, presenting a negative linear effect of the inclusion of SL (P = 0.01; Table 2). In urine, cumulative emissions were 1.44, 0.23 and 0.02 mg m⁻² for 0SL, 50SL, and 100SL, respectively, also presenting a negative linear effect of the inclusion of SL (P = 0.01). The emission factor of N₂O (EFₐ) for each level of inclusion of SL within each type of excreta is described on Table 2.

A linear decrease was observed for cumulative emissions of CH₄ in feces and urine when SL was added (P = 0.03 and P = 0.04, respectively). In feces, a negative cumulative emission, or an uptake of CH₄, was demonstrated by the inclusion of 100% SL (~0.004 mg m⁻²), while in urine, the inclusion of 50% SL presented the least emission (or greatest uptake) in 32 days (~0.013 mg m⁻²). The cumulative emission of CO₂ in feces and urine followed a similar pattern of CH₄, with a linear uptake of CO₂ for the inclusion of SL (P = 0.03) for both excreta types, in which the least emissions were observed for diet 100SL (15.51 and 8.93 mg m⁻² for feces and urine, respectively).

Discussion
In a concomitant study, we demonstrated that a great proportion of CT from SL was bound to protein and fiber in the feces of animals fed 100SL, resulting in lesser apparent total tract digestion of those nutrients than animals consuming grass-only diets. Thus, host enzyme digestion and intestinal absorption of amino acids were inhibited by CT binding ability throughout the gastrointestinal tract, which also contributed to greater concentration of N in feces from animals fed 100SL than feces of animals fed 0SL. Moreover, the greater excretion of N in feces...
may have had a significant contribution from endogenous metabolic N with the addition of CT in the diets18, which is consistent with the studies of Komolong et al.19 and Carulla et al.8. Greater N concentration in urine of animals fed 100SL than 50SL and 0SL can be also attributable to a high CP intake. High CP intake without a concomitant increase in energy supply overloads ruminal microbes, increasing ammonia production in the rumen, thereby, leading to more ammonia being absorbed by the ruminal wall of animals fed 100SL than 0SL20. With that, more ammonia was proportionally being converted to urea and excreted in the urine from animals fed 100SL than those fed the other diets21. Broderick22 states that feeding ruminants exclusively legumes can result in intense urea production, increasing N excretion in the urine, and thus, reducing N retention by the animals. This is because the rate of ammonia production in the rumen exceeds the rate of use by microorganisms23, and if it is not captured as a microbial protein in the rumen, it will undergo the ureagenesis process in the liver.

Beauchemin et al.24 supplemented a 70% forage-based diet with 1 and 2% tannins from quebracho (Schinopsis lorentzii) and reported a decrease in CP digestibility as well as lesser urinary N, as proportion of total N excretion, when CT was included. They also observed an increase in fecal N excretion, as a proportion of total N excretion, with greater inclusion of CT in the diet, which was attributed to endogenous and microbial contributions, and not an actual decrease in urinary N.

The lack of effect of inclusion of SL in the flux of N2O from feces was driven mainly by the presence of organic N in conjunction with the bound-CT5,17. The fluxes of N2O from urine patches are generally greater than fluxes from feces since urea is more rapidly hydrolyzed than organic N. A review21 demonstrated that up to 3.8% of N in urine versus only up to 0.7% of N in feces could be emitted in the form of N2O into the atmosphere. However, in the current study, the lesser cumulative emissions from urine patches than from feces pile are probably because urine was being emitted as another compound, such as volatile ammonia.

The fecal pile functions as an organic fertilizer for the soil and nutrient availability is a key component for soil microbiota. According to the “Hole-in-the-pipe” model proposed by Firestone and Davidson26, the magnitude of

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**Figure 3.** Net flux of CH4 emitted by (a) feces from beef steers fed sericea lespedeza [SL; Lespedeza cuneata (Dum. Cours.) G. Don], where values represent the average of three diets (0, 50 or 100% SL) in two experimental periods, and means followed by different letters are significantly different among sampling days (Ps<sub>day</sub> < 0.001); and (b) urine collected from beef steers fed 0SL, 50SL or 100SL, where asterisk indicates differences among diets within day (Ps<sub>SL*Day</sub> < 0.001). Bars refer to standard deviation. Dashed line represents soil (no excreta) emissions.
of N2O production is mainly a function of the availability of N in the soil. Therefore, the availability of mineral N (NH4+ and NO3−), as well as the availability of labile carbon, promoted greater N2O and CO2 emissions in the first days for all treatments, mainly by activation of microbial populations30. The deposition of fresh excreta triggered denitrification in the initial days, due to greater moisture and N and other nutrients. Moreover, nitrate leaching losses could also have occurred, which, in part, contributes to indirect N2O emissions if nitrate is denitrified within surface waters31,32.

* Figure 4. Net flux of CO2 emitted by (a) feces from beef steers fed 0, 50 or 100% sericea lespedeza [SL; Lespedeza cuneata (Dum. Cours.) G. Don], where values represent the average of three diets (0, 50 or 100% SL) in two experimental periods, and means followed by different letters are significantly different among sampling days (P_day < 0.001); and (b) urine collected from beef steers fed 0SL, 50SL or 100SL, where asterisk indicates differences among diets within day (P_SL*Day < 0.001). Bars refer to standard deviation. Dashed line represents soil (no excreta) emissions.

** Table 1.** Dry matter, nitrogen (N), and condensed tannins (CT) concentration of feces and N concentration composition of urine from beef steers fed three levels of inclusion of sericea lespedeza [SL; Lespedeza cuneata (Dum. Cours.) G. Don], ND not detected. *0SL: 0% inclusion sericea lespedeza hay (100% bermudagrass); 50SL: 50% inclusion sericea lespedeza hay; 100SL: 100% inclusion sericea lespedeza hay (0% bermudagrass hay).
than N2O. This trend supports the hypothesis that more N from urine was being lost as ammonia excreta application), represented by the strong fractionation of ammonia volatilization and the intensification of leaching and consumption by microorganisms. However, CT have been associated with a direct negative effect on methanogens, which might have limited methanogenesis in the excreta of animals fed SL, explaining the linear reduction from 0 to 100% inclusion of the legume in the diets.

In the current study, rainfall events after application of excreta might have reduced emissions of CH4, CO2, and N2O. In our study, the δ15N of N2O was more depleted when emissions were more intense (initial days after excreta deposition), which demonstrates how emissions are more relevant immediately after excreta deposition. According to Franzluebbers and Steiner, temperature, humidity, and soil inorganic N are also determinant of the magnitude of N2O emissions. In the current study, rainfall events after application of excreta might have consequently affected the proportion of water-filled pore space (WFPS), creating anaerobic conditions. Hence, the anaerobiosis and the elevated levels of N and C available in the soil due to the deposition of excreta favored nitrification, nitrifier denitrification, and denitrification, resulting in greater GHG production and emissions. Posteriorly, with elevated temperatures during summer in Florida, a fast drying of the feces and the formation of a superficial crust were observed, which could also have affected the N2O fluxes from feces to decrease from day 5 onward.

Gaseous losses were shown to have a linear relationship with soil δ15N, with large fractionation factors if NO3- is completely consumed in the denitrification reaction. Reported soil and soil-emitted N2O δ15N values typically range between 0 and ~40‰, indicating a wide variation in the depletion of 15N in the gaseous loss pathway. In our study, the δ15N of N2O was more depleted when emissions were more intense (initial days after excreta application), represented by the strong fractionation of ammonia volatilization and the intensification of denitrification process. This trend supports the hypothesis that more N from urine was being lost as ammonia than N2O.

Kool et al. demonstrated that synthetic urine with high hippuric acid concentration reduced N2O emissions by up to 50%. More recently, Zhou et al. attributed the greater reductions in N2O emissions from urine to the addition of tannic acid in the diets of beef cattle. Authors concluded that the increased ratio of hippuric acid-N/urinary N had inhibitory effects in the emissions of N2O, but the effect of the tannic acid was more evident at greater levels of crude protein. Although we did not determine these compounds in the urine, aromatic compounds from CT present in urine from animals fed 100SL had possibly inhibited the nitrification processes in soil to some extent, resulting in lesser emissions of N2O.

Cumulative emissions of all three GHG evaluated were greater for animals fed 0SL than 100SL, regardless of the type of excreta. However, fluxes of CH4 were negative after few days the excreta were applied, indicating an uptake from the soil. Moreover, more than 65% of the total emission of all GHG had been emitted by day 7, which demonstrates how emissions are more relevant immediately after excreta deposition.

Methane emissions from excreta occurs mainly because of the decomposition of feces by methanogenic organisms when in favorable anaerobic conditions. Thus, greater fluxes of CH4 in the first days after application of excreta, regardless of the type, could be observed, which is similar to those observed in the literature in tropical pastures. The methanogenesis was probably favored by the high moisture of the soil in the initial days and tended to decrease with natural decrease of moisture of the excreta, scarcity of rainfall events, and nutrient leaching and consumption by microorganisms. However, CT have been associated with a direct negative effect on methanogens, which might have limited methanogenesis in the excreta of animals fed SL, explaining the linear reduction from 0 to 100% inclusion of the legume in the diets.

In summary, this study showed how diet of the animal, type of excreta, and weather conditions affected GHG fluxes in warm-climate pastures in North Florida. Therefore, feeding beef steers SL was effective in mitigating the emission of N2O, CH4, and CO2 from their excreta for 32 days after deposition in the soil. Furthermore, the EFCH4 of the excreta decreased with the inclusion of tannin-rich legume, but feces and urine should be considered separately due to specific variations in each study conditions (diet, soil, precipitation). Thus, the adoption of specific EFCH4 for countries and regions would avoid overestimation of global emissions from livestock sector. In the current study, more N was added via excreta of animals fed SL hay, reducing N2O emissions compared to excreta of animals fed only grass hay. The δ15N followed a similar pattern of the fluxes of N2O, which was more

### Table 2. Cumulative emissions of greenhouse gases and emission factor of N2O-N from feces and urine of beef steers fed three levels of inclusion of sericea lespedeza [SL; *Lespedeza cuneata* (Dum. Cours.) G. Don].

| Item         | Diets* | 0SL  | 50SL  | 100SL | SEM  | P-valuea |
|--------------|--------|------|-------|-------|------|----------|
| Cumulative emissions, mg N m⁻² |        |      |       |       |      |          |
| **Feces**    |        |      |       |       |      |          |
| N2O          | 2.02   | 1.09 | 0.84  | 0.10  | 0.01 | 0.05     |
| CH4          | 1.98   | 0.04 | 0.004 | 0.06  | 0.03 | 0.90     |
| CO2          | 399.72 | 89.73| 15.51 | 16.4  | 0.03 | 0.18     |
| **Urine**    |        |      |       |       |      |          |
| N2O          | 1.44   | 0.23 | 0.02  | 0.005 | 0.01 | 0.14     |
| CH4          | 0.02   | 0.13 | 0.006 | 0.008 | 0.04 | 0.05     |
| CO2          | 313.74 | 10.81| 8.93  | 24.9  | 0.03 | 0.10     |
| Emission factor of N2O-N, % |        |      |       |       |      |          |
| **Feces**    |        |      |       |       |      |          |
|             | 0.09   | 0.03 | 0.002 | 0.005 | <0.01| 0.25     |
| **Urine**    |        |      |       |       |      |          |
|             | 0.11   | 0.04 | 0.009 | 0.001 | <0.01| 0.13     |

*OSL: 0% inclusion sericea lespedeza hay (100% bermudagrass); 50SL: 50% inclusion sericea lespedeza hay; 100SL: 100% inclusion sericea lespedeza hay (0% bermudagrass hay). BG × SL: bermudagrass versus sericea lespedeza hay inclusion; L: linear effect of SL inclusion; Q: quadratic effect of SL inclusion.
depleted at high release rates due to the activity of soil microorganisms after excreta application. Further research is warranted to understand the mechanisms behind the reduced losses, while studies using additional isotope measurements, such as hydrogen isotopes, would facilitate estimates of emission and atmospheric modeling.

Methods

Animal care. The study followed all procedures approved by the University of Florida Institutional Animal Care and Use Committee (Protocol #201810218) and all methods were carried out in accordance with relevant guidelines and regulations. This manuscript is reported in accordance with ARRIVE guidelines.

Experimental site. The experiment was conducted at North Florida Research and Education Center (NFREC), from University of Florida, located in Marianna, FL (30°52’ N, 85°11’ W, 35 m a.s.l.) in a pasture of ‘Pensacola’ bahiagrass (Paspalum notatum Flüggé). The soil at the experimental site is classified as an Orangeburg loamy sand (fine-loamy-kaolinitic, thermic Typic Kandiudults), with an average pH of 6.5. Average Mehlich-I extractable P, K, Mg, and Ca concentrations were 13, 45, 31, and 245 mg kg−1, respectively. Soil organic matter was 6.3 g kg−1, and the estimated cation-exchange capacity was 2.8 meq 100 g−1. The study was carried out for two experimental periods of 32 days each, separated by a 15-day interval (Period 1: from 06/08/2018 to 07/10/2018; Period 2: from 07/25/2018 to 08/27/2018). The average, maximum and minimum temperatures and rainfall for the experimental periods are represented in Fig. 5.

Donor animals and experimental diets. Fifteen Brahman × Angus crossbred steers [Period 1: 324 ± 26 kg initial body weight (BW); Period 2: 336 ± 30 kg BW] were randomly distributed into three experimental diets: 0, 50, or 100% (as fed) inclusion of ‘AU Grazer’ sericea lespedeza hay [SL; Lespedeza cuneata (Dum. Cours.) G. Don] into ‘Tifton 85’ bermudagrass hay (BG; Cynodon spp.) diets (Table 3) and used as donors of excreta (urine and feces). Steers were fed for 21 days for two feeding periods in a concomitant study17 and excreta used in the current study was collected in the last two experimental days of each. All methods were carried out in accordance with relevant guidelines and regulations.

Static chamber technique. Emissions of GHG were evaluated using the static chamber (non-steady state) technique44. Chambers were circular with 30 cm radius and made of a base and a lid, both built out of PVC47. The lids were wrapped with reflective tape to provide insulation and a rubber septum was added for gas sampling48. The base was fitted with a 10-cm length copper venting tube to ensure adequate air pressure inside the chamber during measurements49,50. Lids and bases were kept closed for gas sampling by fitting a bicycle tire inner tube that tightly sealed the parts together.

Figure 5. Marianna (FL) station of the Florida Automated Weather Network (FAWN) data of (a) weekly rainfall and temperature data from two experimental periods and (b) accumulated monthly rainfall (mm) and average temperature (°C). Period 1: from 06/08/2018 to 07/10/2018; Period 2: from 07/25/2018 to 08/27/2018.
Bases of chambers were installed in the non-grazed pasture of bahiagrass two weeks prior to excreta application to avoid any effect of soil disturbance on GHG emissions. Bases were installed at 8-cm depth, with 5 cm extending above ground level. Depth for installation was determined based on Clough et al. Chamber tops were 22 cm height, which when summed with 5 cm of base totaled 27 cm, in agreement with the indication of ≥ 40 cm of chamber height per hour of deployment. New bases were installed in a near location of the same pasture for the second experimental period, also two weeks prior to starting new gas sampling.

### Treatments, excreta deposition, and gas sampling.

Treatments applied to the chambers consisted of either feces or urine within one of the three levels of inclusion of SL hay fed to the beef steers and were distributed as a complete randomized block design. Urine and feces were collected directly from each animal by spontaneous or stimulated urinations and defecations and applied at a rate of 2 L of urine and 2 kg of feces, as typical amounts excreted by cattle for the area of the chamber. To obtain quantities required of excreta, sampling occurred twice a day (700 and 1500 h) and samples were kept refrigerated at 4 °C until next morning (day 0 of gas sampling). Samples of each excreta type were composited across all five animals within each SL diet resulting in three final subsamples (urine and feces from each of three SL diets). Excreta samples were kept at room temperature 2 h prior to application to the chambers and their chemical composition is described on Table 1.

Application of excreta to the chambers was made one time in each experimental period on the soil surface inside the area determined by the base of the chamber. Grass inside the chamber area was cut at ground level before each sampling day, when appropriate. Gas sampling occurred between 0900 and 1100 h, when temperature is considered more representative of the daily average on days 0, 1, 3, 5, 7, 14, 18, 25, and 32 after excreta application for both experimental periods. One subsample was taken per deployment time per chamber, separated by 15-min intervals (T₀, T₁₅, and T₃₀). At T₀ a sample was collected from the area directly above the soil surface. Immediately thereafter, chambers were tightly closed by fitting the lid to the base with the bicycle inner tube, followed by the next sample deployment times. All samples were collected with the use of a 60-mL syringe and immediately flushed into a pre-vacuumed 30-mL glass vial. The vial was equipped with a butyl rubber stopper and sealed with an aluminum septum. Samples were analyzed immediately after finishing each experimental period.

Gas sample analyses were conducted using a gas chromatograph (Trace 1310 Gas Chromatograph, Thermo Scientific, Waltham, MA). For N₂O, an electron capture detector (350 °C) and a capillary column (J&W GC packed column in stainless steel tubing, length 6.56 ft (2 M), 1/8 in. OD, 2 mm ID, Hayesep D packing, mesh size 80/100, pre-conditioned, Agilent Technologies) were used. Methane was analyzed using a flame ionization detector (250 °C) and a capillary column (J&W PoraBOND Q GC Column, Agilent Technologies). For CO₂, a thermal conductivity detector (200 °C) and capillary column (J&W GC packed column in stainless steel tubing, length 7 ft (2.13 M), 1/8 in. OD, 2 mm ID, Haysep N packing, mesh size 60/80, pre-conditioned, Agilent Technologies) were used. Temperature of the injector and columns were 80 and 200 °C, respectively.

### Calculations and δ¹⁵N-N₂O composition.

The hourly gases fluxes (mg of N₂O or CH₄ or CO₂ per m⁻² h⁻¹) were calculated according to Cardoso et al.:

\[
F_{\text{GHG}} = \left(\frac{\delta C}{\delta t}\right) \times \left(\frac{V}{A}\right) \times \left(\frac{M}{V_m}\right),
\]

where \(\delta C/\delta t\) is the change in gas concentration in the chamber during the deployment time; V and A are the chamber volume and soil area covered by the chamber, respectively; M is the molecular weight of the gas; and Vm is the molecular volume of gas. The Vm parameter was corrected to the standard conditions of temperature and pressure as Vm = 0.02241 × (273.15 + Tc/273.15) × p0/p1, where 0.02241 is the molar volume (m³). Tc is the

### Table 3. Chemical composition (DM basis) of experimental diets containing sericea lespedeza [SL; Lespedeza cuneata (Dum. Cours.) G. Don] hay fed to beef steers donors of excreta (adapted from van Cleef et al. 17).

| Item                   | Dietary treatmentsa | OSLa | 50SL | 100SL |
|------------------------|---------------------|------|------|-------|
| DM, g/kg               |                     | 898  | 922  | 913   |
| Nutrient composition, g kg⁻¹ DM |                     |      |      |       |
| OM                     |                     | 829  | 867  | 857   |
| CP                     |                     | 85   | 108  | 161   |
| NDF                    |                     | 747  | 389  | 377   |
| ADF                    |                     | 345  | 323  | 258   |
| Condensed tannins, g kg⁻¹ DM |                     |      |      |       |
| Bound                  |                     | ND   | 23.8 | 53.5  |
| Unbound                |                     | ND   | 12.5 | 28.8  |
| Total                  |                     | ND   | 36.3 | 82.3  |
chamber headspace temperature at sampling time (°C), p0 is the air pressure at sea level, and p1 is the local pressure calculated using the barometric equation. The minimal detectable flux was 0.012 ppb min⁻¹ for N₂O, 0.004 ppm min⁻¹ for CH₄, and 1.40 ppm min⁻¹ for CO₂.

Daily N₂O, CO₂, and CH₄ emissions were calculated by multiplying the fluxes by 24 h and cumulative emissions were estimated by integrating the fluxes over each day (area under the curve) and averaged per period. The fraction of N applied in the excreta lost as N₂O, named as emission factor (EF), was calculated according to the equation:

$$\text{EF}_{\text{N}_2\text{O}}(\%) = \left(\frac{\text{N}_2\text{O} - \text{N}_{\text{emitted}}}{\text{N}_2\text{O} - \text{N}_{\text{blank}}}\right) \times 100,$$

where \(\text{EF}_{\text{N}_2\text{O}}\) is the emission factor of N₂O; \(\text{N}_2\text{O}-\text{N}_{\text{emitted}}\) is the cumulative N₂O-N emissions from the chamber with excreta (mg m⁻²); \(\text{N}_2\text{O}-\text{N}_{\text{blank}}\) is the cumulative N₂O-N emissions from the blanks (chamber with no excreta deposited; mg m⁻²); and \(\text{N}_{\text{applied}}\) is the urine or feces N application rate (mg m⁻²).

A subsample (12 ml) of the collected gas was transferred to an isotope ratio mass spectrometer (IsoPrime 100, IsoPrime, Manchester, UK) for ¹⁵N analysis, using He (2 ml min⁻¹) as a carrier. The isotope ratio for ¹⁵N/¹⁴N was calculated as:

$$\delta^{15}N = \left(\frac{^{15}N/^{14}N_{\text{sample}} - ^{15}N/^{14}N_{\text{reference}}}{^{15}N/^{14}N_{\text{reference}} \times 1000}\right),$$

where \(\delta^{15}N\) is the N isotope ratio of the sample relative to atmospheric nitrogen. \(^{15}N/^{14}N_{\text{Sample}}\) is the N isotope ratio of the sample, and \(^{15}N/^{14}N_{\text{reference}}\) is the N isotope ratio of atmospheric N (standard). The stable isotopic composition of nitrogen was reported using the conventional delta per mill notation. \(\delta^{15}N\) values are expressed relative to the international standard (AIR-N₂).

Statistical analyses. The experiment was analyzed as a completely randomized block design, with feces and urine data computed separately due to differences in the magnitude of responses. There were three replicates (chamber) of each treatment, and day was considered the repeated measurement for all variables. Glimmix procedure of SAS (SAS Inst., Inc., Cary, NC, version 9.4) was used, in which the chamber was considered the experimental unit. Graphs were drawn using Microsoft Excel (version 16.61). Normality of distribution and homogeneity of variances were evaluated using the Univariate procedure of SAS. Covariance structures were based upon the smallest Akaike Information Criterion value. The model included the fixed effect of level of SL composition of nitrogen was reported using the conventional delta per mill notation. \(\delta^{15}N\) values are expressed relative to the international standard (AIR-N₂).

Data availability
All data generated or analyzed during this study are included in the article.

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Author contributions
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Competing interests
The authors declare no competing interests.

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