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## Different methods and times to estimate heat production in sheep fed with sunflower meal

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**ABSTRACT**: The objective of this study was to assess the oxygen pulse and heart rate method ( $O_2P$ -HR) using a respiration chamber at different measurement times for estimate the heat production (HP) of lambs fed increasing levels of sunflower meal in their diet. Twenty-four lambs were assigned to four experimental diets (0, 100, 200, and 300 g of sunflower meal kg<sup>-1</sup> DM). Heat production was estimated using the  $O_2P$ -HR (HP<sub>02P</sub>) method and a respirometry chamber (HP<sub>RC</sub>). Measurements were obtained by simultaneously measuring heart rate (HR) and oxygen consumption over 3, 6, 9, 12, 15, 18, 21 and 24 h. A flow-through respirometry chamber for small ruminants was used to determine oxygen consumption (VO<sub>2</sub>) and carbon dioxide and methane production. Data on dietary treatment, measurement times and their interactions were analyzed as repeated measures using mixed model procedures and Restricted Maximum Likelihood (REML) estimation. The Pearson's correlation coefficient was used to compare techniques. There was no effect of the different levels of sunflower meal inclusion on VO<sub>2</sub> and heat production. The HP<sub>02P</sub> (126.16 kcal/ BW<sup>0.75</sup>/day) was 2% higher than that of the HP<sub>RC</sub> (124.61 kcal/ BW<sup>0.75</sup>/day), and the correlation coefficients was 0.628. The coefficient of variation was greater for the HP<sub>02P</sub> (21.33%) than for HP<sub>RC</sub> (11.44%). HR (beats/min), VO<sub>2</sub> (mL/min/BW<sup>0.75</sup>) and oz<sub>2</sub>P-HR (mL/beat) required measurement times of 24, 15 and 9 hours, respectively. A measurement time of 24 h was necessary to ensure a more accurate estimate of the heat production using the O<sub>2</sub>P-HR method. **Key words**: bioenergetic, energy requirements, indirect calorimetry, lamb, ovine.

# Diferentes métodos e tempos de medição para estimar a produção de calor em ovinos alimentados com farelo de girasssol

**RESUMO**: O objetivo com este estudo foi avaliar o método do pulso de oxigênio ( $O_2P$ -FC) usando câmara respirométrica em diferentes tempos de medição para estimar a produção de calor de cordeiros alimentados com níveis crescentes de farelo de girassol na dieta. Vinte e quatro cordeiros foram distribuídos em quatro dietas experimentais (0,10, 20 e 30% de farelo de girassol). A produção de calor foi estimada pelo método de  $O_2P$ -FC ( $PC_{O2P}$ ) e por câmara respirométrica ( $PC_{CR}$ ). As estimativas foram obtidas medindo-se simultaneamente a frequência cardíaca (FC) e o consumo de oxigênio ( $VO_2$ ) durante 3, 6, 9, 12, 15, 18, 21 e 24 horas. Uma câmara de respirométrica para pequenos ruminantes foi usada para determinar o  $VO_2$  e a produção de dióxido de carbono e metano. Os dados referentes a dieta experimental, tempos de medição e suas interações foram analisados como medidas repetidas usando os procedimentos de modelo misto e estimativa de máxima verossimilhança restrita. A correlação de pearson foi usada para comparar as duas técnicas de estimativa da produção de calor. Não houve efeito dos diferentes níveis de inclusão de farelo de girassol sobre o consumo de oxigênio e produção de calor dos animais. A PC<sub>02P</sub> (126,16 kcal/ PV<sup>0.75</sup>/dia) foi 2% maior que a PC<sub>CR</sub> (124,61 kcal/ PV<sup>0.75</sup>/dia), e o coeficiente de correlação foi de 62,8%. O coeficiente de variação foi maior para PC<sub>O2P</sub> (21,33%) comparado com PC<sub>CR</sub> (11,44%). A FC (batimentos/min), VO<sub>2</sub> (mL/min/PC<sup>0.75</sup>) e o O<sub>2</sub>P-FC (mL/batimento) requerem tempos de medição de 24, 15 e 9 horas, respetivamente. É necessário a mensuração por 24 horas para garantir uma estimativa mais precisa da produção de calor usando o método de O<sub>2</sub>P-FC.

Palavras-chave: bioenergética, calorimetria indireta, cordeiros, exigências de energia, ovinos.

#### **1 INTRODUCTION**

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3 Energy is the primary nutrient limiting 4 ruminants. It is derived from the oxidation of dietary 5 nutrients and is essential for maintenance of vital 6 processes. This nutrient is dissipated by animals during 7 ingestion and metabolism of food: first, energy is consecutively lost in feces, urine and fermentative gases, and subsequently lost as heat increases. The remaining energy is primarily directed towards maintenance (approximately 70% of the net energy available) and production processes (TEIXEIRA et al., 2017).

The standard method for measuring energy expenditure in ruminants involves the use of open-

1 circuit respirometry chambers. In this method, the 2 products resulting from the animal's metabolism, 3 such as gas exchanges with the environment (oxygen consumption, carbon dioxide and methane 4 5 production), combined with urinary nitrogen excretion, are quantified (SILVA, 2011; OSS et 6 7 al., 2016). A respirometry chamber is an accurate technique, however it is used under laboratory 8 9 conditions and is extremely expensive. It also requires 10 significant expertise and infrastructure, wich makes it 11 impractical for small rural properties (RODRIGUEZ 12 et al., 2007; MACHADO et al., 2016).

13 Measuring the heat production of animals 14 can provide insights into how efficiently they utilize 15 the nutrients in their diet, thereby helping optimize feed 16 efficiency, since heat production is closely linked to the 17 metabolic processes and energy utilization of animals.

Researchers are seeking to estimate heat 18 19 production in ruminants by using heart rate adjusted 20 for oxygen consumption per beat as there is a linear 21 relationship between heart rate (HR) and oxygen 22 consumption  $(VO_2)$  in homeothermic animals, 23 thereby indicating that it is possible to estimate heat production through HR measurements (TALMON 24 25 et al., 2023). The primary goal is to improve and 26 develop techniques capable of measuring the energy 27 requirements of animals in a shorter time frame, with cheaper equipment, and without changing the 28 29 behavior and normal conditions of animal husbandry 30 (BROSH, 2007; CHAVES et al., 2015).

31 The O<sub>2</sub>P-HR technique can be used as an 32 alternative method to determine heat production. 33 However, there are some problems associated with 34 the ideal time to measure oxygen consumption, heart rate, and O<sub>2</sub>P-HR, especially considering the 35 intraday changes that interact directly with animals. 36 37 According to OSS et al. (2016), O<sub>2</sub>P-HR is an 38 alternative technique, but has a greater between-39 animal coefficient of variation, which has a negative 40 effect on the power of the experiments. Further 41 studies should be performed to investigate ways to minimize the errors associated with the O<sub>2</sub>P-HR 42 43 method to increase the precision and statistical power 44 of experiments using this technique.

The objective of this study was to evaluate
the O<sub>2</sub>P-HR method at different measurement times
to estimate heat production in crossbred lambs fed
diets containing increasing levels of sunflower meal.

#### 50 MATERIALS AND METHODS

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52 Twenty-four crossbred (Santa Inês x 53 Dorper) intact male lambs with a mean age of 4 months were arranged into three blocks, four treatments, and two replicates per block using a randomized block design. A 10-day adaptation period was allowed before the data collection.

The lambs received four isoproteic experimental diets formulated according to the NRC (2007) recommendations for lambs on maintenance levels. The diets containing a roughage:concentrate ratio of 40:60 on a dry matter basis (DM). Corn silage was supplied as the roughage source and the concentrate was formulated by replacing soybean meal with increasing levels of sunflower meal (0, 100, 200 and 300 g kg<sup>-1</sup> DM) (Table 1). The diet was provided in two daily meals at 8 a.m. and 4 p.m.

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The chemical composition of the diets and orts was determined by analyzing the dry matter (DM), ash, crude protein (CP), neutral detergent fiber (NDF), acid detergent fiber (ADF), ether extract (EE), non-fibrous carbohydrates (NFC) and total carbohydrate (TC) content according to the procedures of INCT-CA (DETMANN et al., 2012).

Before starting the experiment, the lambs were weighed, drenched and vaccinated against clostridial diseases. During the experiment, lambs were housed in individual metabolic cages provided with feed and water troughs, which allowed the collection of urine and fecal samples. After the adaptation period DM intake was measured, and urine was collected for nitrogen determination for five days. Heat production was estimated by using a respirometry chamber after adaptation to the diet.

The  $O_2$  consumption (VO<sub>2</sub>) and CO<sub>2</sub> production data were recorded using a Sable System (Sable Systems International, Las Vegas, NV, USA). The lambs were individually placed in a respirometry chamber for 24 h and the same dietary treatment offered during the adaptation period was administered to each lamb once in the morning.

Ambient air flowed through the chamber at a controlled flow rate based on lamb's weight (0.6 liters/ kg of body weight/minute), and it was mixed with the exhaled air. Samples were taken every 5 min for 24 h to determine  $O_2$ ,  $CO_2$ , and  $CH_4$  concentrations. All data were recorded using an automated data acquisition program (Expedata; Sable Systems International).

The maximum allowable concentration of  $CO_2$  in the chamber was 1.0%. Oxygen consumption and  $CO_2$  production were calculated by comparing the composition and volume of the air that flowed through the respirometry chamber with the air released. The temperature was kept at 22 °C by using an air conditioner placed inside the chamber to provide thermal comfort to lambs.

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Item (g kg <sup>-1</sup> )	Sunflower meal level (g kg <sup>-1</sup> DM)							
	0	100	200	300				
Corn silage	400	400	400	400				
Soybean meal	264	196	118	18				
Corn grain	315	281	256	261.6				
Sunflower meal	0	100	200	300				
Vitamin-Mineral Premix <sup>1</sup>	10.5	15.0	22.0	36.6				
Dicalcium phosphate	10.5	8.0	4.0	0				
Nutrients								
DM (g/kg <sup>-1</sup> NM)	643.4	641.7	653.3	658.8				
MM (g/ kg <sup>-1</sup> DM)	45.0	43.2	45.7	46.1				
CP (g/ kg <sup>-1</sup> DM)	207.9	203.8	195.6	189.5				
NDF(g/ kg <sup>-1</sup> DM)	337.9	364.5	391.2	419.8				
ADF(g/ kg <sup>-1</sup> DM)	176.3	212.5	227.2	302.8				
EE (g/ kg <sup>-1</sup> DM)	65.5	62.9	60.9	60.1				
NFC(g/ kg <sup>-1</sup> DM)	343.7	325.6	306.6	284.5				
TC (g/ kg <sup>-1</sup> DM)	681.6	690.1	697.8	704.3				

Table 1 - Nutritional composition of the experimental diets.

DM = Dry matter, NM = Natural Matter, CP = Crude Protein, NDF = Neutral detergent Fiber, ADF = Acid detergent fiber, EE = Ether extract, NFC = Non-fibrous carbohydrate, TC = Total carbohydrate.

<sup>1</sup>Composition of Vitamin-Mineral Premix: Calcium (Max.) 150 g, Calcium (Min.) 130 g, Phosphorus (Min.) 65 g, Sodium (Min.) 130 g, Fluorine (Max.) 50 mg, Sulfur (Min.) 12 g, Magnesium (Min.) 10 g, Iron (Min.) 1000 mg, Manganese (Min.) 3000 mg, Cobalt (Min.) 80 mg, Zinc (Min.) 5000 mg, Iodine (Min.) 60 mg, Selenium (Min) 10 mg, Vitamin A (Min.) 50000 IU, Vitamin E (Min.) 312 IU.

1 Heat production was estimated using the 2 respirometry chamber technique  $(HP_{RC})$  according to

- 3 Brouwer's equation (1965) as follows:
- 4  $HP_{RC}(Kj) = 16.18 \times VO_2(L) + 5.02 \times VCO_2(L) 5.88$
- 5 x UN (g) -2.17 x VCH<sub>4</sub>(L)
- $6 \text{ HP}_{\text{RC}} = \text{Estimation of heat production using the respi-}$
- 7 rometry chamber technique
- 8  $VO_2 = Oxygen consumption$
- 9  $VCO_2 = Carbon dioxide production$
- 10 UN = Urinary nitrogen
- 11  $VCH_4 =$  Methane production

12 Estimation of heat production using the  $O_2$  pulse methodology (HP<sub>02P</sub>) was based on a pro-13 tocol adapted from BROSH et al. (1998). After 14 15 the adaptation period, the lambs were monitored 16 for four days to record the mean heart rate using 17 a POLAR<sup>®</sup> RS800 transmitter. The transmitters were attached to the girth of the lambs using elas-18 19 tic strips. Data were recorded at 60 s intervals and 20 subsequently transferred to a computer using an 21 infrared sensor.

22 After determining the mean heart rate 23 (HR during the four days of measurement), data on 24 heartbeat (HR-RC) and oxygen consumption (VO<sub>2</sub>) 25 were collected simultaneously for 24 h using a 26 respirometry chamber, as described above. These data were used to calibrate the  $O_2$  volume per heartbeat. The oxygen pulse and heart rate were calculated as  $VO_2$  per heartbeat.

Daily heat production was obtained by multiplying the total  $O_2$  consumption by the constant 4.89 kcal/L of  $O_2$  (NICOL & YOUNG, 1990). The results were expressed as metabolic weight (kcal/kg BW<sup>0.75</sup>/day). Heat production was estimated using the following equation:

 $HP02P: \frac{kcal}{day \times kg BW0.75} = (HR - RC X 2 O2P X 4.89) X 1440 / (kg BW 0.75)$ 

 $HP_{O2P}$  = Estimation of heat production using the oxygen pulse and heart rate method

HR-RC: Mean heartbeat (beat/min)

O<sub>2</sub>P: Oxygen consumption per heartbeat (L/beat).

#### Data analysis

The dietary treatments, measurement times and their interactions were analyzed as repeated measures (each treatment was analyzed at eight measurement times: 3, 6, 9, 12, 15, 18, 21 and 24 h) because the observations were interdependent. Data were analyzed using the Proc MIXED procedure in SAS (SAS 9.0 Inst. Inc.) and Restricted Maximum Likelihood (REML) estimation according to the following model:

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1  $yijt = \mu + \alpha i + dj(i) + \gamma t + (\alpha \gamma)it + (b + \beta j) + eijt$ 

2 Where, yijt = the expected outcome 3 for the dependent variable Y observed at the measurement time t for the lamb j fed the diet i;  $\mu$  is 4 5 the overall mean;  $\alpha i$  is the fixed effect of diet; di(i)is the random effect of lamb *j* nested within diet 6 7 *i*;  $\gamma t$  is the fixed effect of measurement time; ( $\alpha \gamma$ ) it is the interaction between diet and measurement 8 9 time; b is the regression coefficient;  $\beta i$  is the slope 10 deviation (diet *i*) of the regression coefficient *b*; eijt 11 is the random error associated with lamb *j* fed diet *i* 12 at the measurement time t, eijt ~ NID ((0,  $\sigma^{e}_{2}$ ) (data 13 is approximately normally distributed with mean of 0 and variance of  $\sigma_{2}^{e}$ ; and the values of dj(i) and eijt 14 are assumed to be independent. 15

16 Five variance-covariance matrix structures 17 were tested as follows: variance components (VC variances are equal and observations are independent, 18 i.e., there is no correlation between observations 19 over time); compound symmetry (CS - equality of 2021 variances and covariances); first-order autoregressive model (AR (1) - equality of variances and covariances 22 23 with higher correlation between adjacent measures); first-order ante-dependence (ANTE (1) - the 24 25 magnitude of the covariance depends on the values 26 of both correlation and standard deviations associated 27 with them); unstructured (UN - each variance and 28 covariance is estimated exclusively from the data) 29 (SAS, 2004). The best model for each set of variables was selected based on the lowest corrected Akaike 30 31 Information Criterion (AICc) value. 32 The variance-covariance matrix structure of the best fit for the measurement time 33 34 was selected based on the lowest corrected Akaike Information Criterion (AICc) value (LITTELL et 35 al., 2006). The ANTE (1) model provided the best 36

fit for HR, O<sub>2</sub>P (mL/beat/BW<sup>0.75</sup>), HPO<sub>2</sub>P (kcal/day)

and HPO<sub>2</sub>P (kcal/day/BW<sup>0.75</sup>), thereby modeling

the covariance structure and thus generating valid tests. The AR (1) model provided the best fit for the VO<sub>2</sub> (L/day), VO<sub>2</sub> (mL/min/BW<sup>0.75</sup>) and O<sub>2</sub>P (L/beat), whereas the ANTE (1) model did not converge. The UN model did not converge for VO<sub>2</sub> (L/day) or VO<sub>2</sub> (mL/min/BW<sup>0.75</sup>). For the other parameters, problems were encountered when the Hessian matrix was applied, which demonstrates that the UN structure was inappropriate.

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After defining the best model for each set of variables, the result of the fixed effect analysis (measurement time) was used as a criterion to test the significance of the treatment effect ( $\alpha$ =0.05). The parameters were subjected to regression analysis (PROC REG) using SAS (2004) when the diet was significant. Differences between groups means (each measurement time vs 24 h) were determined by calculating the minimum significant difference for p = 0.05 using the Tukey's test' when measurement time was significant. To express the accuracy and repeatability of the test, the coefficient of variation was calculated using the PROC UNIVARIATE procedure (SAS, 2004). Pearson's correlation was used to compare techniques using PROC CORR (SAS, 2004). Statistical significance was set at P < 0.05.

## **RESULTS AND DISCUSSION**

The inclusion of sunflower meal did not 30 change (P > 0.05) the DM intake, as the animals were 31 fed at a maintenance level, with averages of 612 g/ 32 day and 50.48 g/BW<sup>0.75</sup>/day. There was no effect on 33 CP intake owing to the lack an effect on DM intake 34 and the isonitrogenous profile of the diets (Table 2). 35 However, there was a significant difference (P < 0.05) 36 in NDF intake among the treatments, with a linearly 37 increasing effect observed. This behavior can be 38

Table 2 - Means, coefficient of variation (CV) for dry matter intake (DMI), crude protein intake (CPI) and neutral detergent fiber intake (NDFI) of lambs fed with different levels of sunflower meal inclusion.

Variables	Sunflower meal level (g kg <sup>-1</sup> DM)				CV	PP		
	0	100	200	300	(%)	Linear	Quadratic	
DMI (g/day)	637.63	596.57	601.7	613.80	10.70	0.593	0.5440	
DMI (g/ BW <sup>0.75</sup> /day)	50.94	50.16	50.28	50.54	2.63	0.6576	0.5850	
CPI (g/ BW <sup>0.75</sup> /day)	10.9	9.85	10.04	10.04	2.72	0.8721	0.5929	
NDFI (g/ BW <sup>0.75</sup> /day) <sup>1</sup>	17.49	19.58	22.47	25.14	14.17	<.0001	<.0001	

 $^{1}y = 17.29 + 0.0258x; R^{2} = 0.967.$ 

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1 explained by the increased fiber concentration in the 2 diets resulting from the inclusion of SFM.

3 The difference in NDF intake was not 4 sufficient to change the VO<sub>2</sub> or heat production 5 (P > 0.05). This behavior was possibly due to the 6 animals' energy intake being close to maintenance 7 levels. Typically, animals with higher digestible 8 energy intake consume more oxygen during 9 metabolic processes.

10 The HR-RC, normal HR, VO<sub>2</sub>, HP<sub>02P</sub>, and 11 HP<sub>RC</sub> did not differ bamong sunflower meal inclusion 12 levels (P > 0.05), possibly because the lambs were fed 13 near maintenance. Moreover, the interaction between 14 SFM level and time was not significant (P > 0.05).

15 The HR of lambs during the four days of 16 measurement varied by only 2.6% in comparison with 17 the HR-RC (Table 3), wich suggests the absence of 18 stress or lack of exercise during the evaluation period. 19 LANDAU et al. (2006) observed similar results for 20 daily HR and HR-RC (81.1  $\pm$  5.1 and 79.2  $\pm$  5.1, 21 respectively). According to the authors, this response 22 was associated with a lower environmental effect on 23 grazing during the measurement of the O<sub>2</sub> pulse.

24 The  $VO_2$  (Table 3) was similar to that 25 reported by MACHADO et al. (2015) when evaluating 26 heat production in sheep fed sorghum silages at 27 different maturation stages (18.37 mL/min/kg BW<sup>0.75</sup>). 28 The similarity in the results is likely related to feeding 29 near maintenance and absence of stress during the 30 execution of both studies. This is also corroborated by 31 the heart rate data as variations of less than 20% were 32 observed between normal heart rates.

33 The O<sub>2</sub>P values (Table 3) corroborate those reported by ARIELI et al. (2002) (0.250 mL/beat/kg/ 34 BW<sup>0.75</sup>) in sheep fed high or low energy diets (75% 35 and 25% concentrate, respectively). Under conditions 36 37 where animals are not subjected to stress, physical

activity or if the variation in heart rate is less than 20%, O<sub>2</sub>P remains constant, and the data are reliable.

The coefficient of variation was greater for  $HP_{O2P}$  (21.33%) than for  $HP_{RC}$  (11.44%). The repeatability of individual animals over time needs to be high to reliably detect the differences in the heat production of animals through respiration trials. OSS et al. (2016) compared the O<sub>2</sub>P method with measurements using a respirometry chamber in crossbred steers (Holstein × Gyr), which was also confirmed by a greater between-animal coefficient of variation (16.6%) compared to RC (7.7%). According to the authors, the O<sub>2</sub>P-HR method had a higher coefficient of variation, and the sample size (n) must be increased to determine the differences in HP between treatments more accurately.

Despite the differences in the coefficients of variation, the correlation between  $HP_{O2P}$  and  $HP_{RC}$ was 0.628 (Figure 1), thereby validating the efficiency of the O<sub>2</sub>P-HR method in predicting heat production. The  $HP_{O2P}$  was 2% higher than the  $HP_{RC}$ . In a study evaluating the efficiency of the O,P-HR method as a tool for determining energy expenditure in sheep, ARIELI et al. (2002) reported that heat production using the O<sub>2</sub>P-HR technique was 6.7% higher than that using the comparative slaughter method.

Table 4 presents the differences between the overall means observed at each measurement time 28 and after 24 h. A significant difference was detected 29 between measurement times of up to 6 h vs. 24 h for all variables studied. This response is probably due to the initial adjustment phase of gas collection, which is essential for equilibrium between gas production and consumption inside the respirometry chamber.

There was an increase in HR and VO, during the first hours, followed by a gradual reduction after feeding. This response is likely related to

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Table 3 - Mean, standard deviation, minimum and maximum of heart rate, oxygen consumption and heat production of sheep.

Item	Mean	SD	Min.	Max.
HR <sup>1</sup> , bpm	79.52	9.28	65.70	96.80
HR-RC <sup>2</sup> , bpm <sup>3</sup>	81.65	11.50	61.84	105.71
VO2 <sup>4</sup> , ml/min/kg BW <sup>0.75</sup>	18.09	2.52	12.21	29.42
$O_2 P^5$ , ml /beat/kg/ BW <sup>0.75</sup>	0.225	0.039	0.143	0.287
HP <sub>02P</sub> <sup>6</sup> , kcal/day/kg BW <sup>0.75</sup>	126.16	26.91	68.72	164.92
$\mathrm{HP_{RC}}^{7}$ , kcal/day/kg BW <sup>0.75</sup>	124.61	14.26	95.94	149.03

<sup>1</sup>Mean heart rate during the four days of measurement. <sup>2</sup>Mean heart rate collected for 24 hours using a respirometry chamber. <sup>3</sup>Beat per minute. <sup>4</sup>Oxygen consumption. <sup>5</sup>Oxygen consumption per heart beat. <sup>6</sup>Heat production using the O<sub>2</sub> pulse methodology. <sup>7</sup>Heat production using the respirometry chamber methodology.

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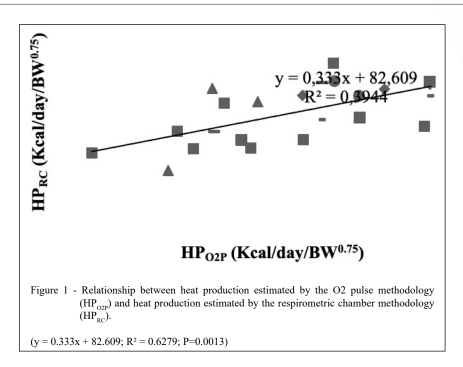
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1 reduced stress levels after the initial period and a more extended post-feeding period, as the diet was provided 2 3 once a day at the beginning of the measurement. The 4 feeding time and the physical activities of chewing 5 and swallowing were the leading causes of the 6 increase in HR and VO<sub>2</sub> during the first few hours, as they showed a gradual reduction after feeding. 7 According to TALMON et al. (2023) eating was the 8

9 activity that most increased HP, VO<sub>2</sub>, and HR.

10 Measurement time had a significant effect

11 on HR (Table 4) throughout the 24 h period (P < 0.0001). Nevertheless, the variations observed after the 9 h period were lower than 15%. According to BROSH (2007), variations in a normal heartbeat are acceptable for the determination of O<sub>2</sub>P, thereby ensuring the reliability of our database.

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Based on the statistical analysis, we observed that HR measurements should be performed for 24 h for more complete data collection, avoiding intraday variations such as feeding time and diet 10 quality (metabolic activity increases during digestion and absorption), lower heart rate at night (when 11

Table 4 - Variation in heart rate, oxygen consumption, oxygen volume per heart beat, and heat production using the O2 pulse methodology in sheep (n = 23), expressed as the difference between the mean measurement at each time studied and that obtained during 24 hours.

			Meas	urement time	;			
Variable	3h	6h	9h	12h	15h	18h	21h	24h
HR, beat/min	26.52***	$17.08^{***}$	12.88***	8.34***	5.17***	2.64***	$0.97^{***}$	$81.65\pm2.40$
VO2, l/day	$5.40^{***}$	4.13**	3.26**	$2.11^{*}$	1.02	-0.16	-0.04	$30.53 \pm 8.92$
VO <sub>2</sub> , ml/min/BW <sup>0.75</sup>	3.19***	2.43**	$1.94^{*}$	1.24*	0.60	0.06	-0.02	$18.09\pm0.53$
O <sub>2</sub> P, ml/beat	-0.30**	$-0.20^{*}$	-0.14	-0.10	-0.09	-0.09	-0.04	2.6±0.11
O <sub>2</sub> P, ml/beat/BW <sup>0.75</sup>	-0.03*	-0.02*	-0.01*	$-0.008^{*}$	$-0.007^{*}$	$-0.007^{*}$	$-0.002^{*}$	$0.225 {\pm} 0.008$
HP <sub>02P</sub> , Kcal/day	$-174.9^{*}$	-107.3*	-79.51*	$-55.98^{*}$	-52.63*	-52.63*	$-20.48^{*}$	$1485.2 \pm 74.32$
HP <sub>O2P</sub> ,Kcal/day/BW <sup>0.75</sup>	-14.51*	-8.75*	-6.48*	-4.60*	-4.23*	-4.23*	-1.63*	126.17±5.61

 $^{***}P < 0.0001;$   $^{**}P < 0.001;$   $^{*}P < 0.05$ . HR=Mean heart rate during the four days of measurement. VO<sub>2</sub>=Oxygen consumption. O2P= Oxygen consumption. per heart beat. HP<sub>02P</sub>=Heat production using the O<sub>2</sub> pulse methodology.

1 animals are at rest) and excitement resulting from 2 the presence of people. BARKAI et al. (2002), 3 ARIELI et al. (2002), AHARONI et al. (2003), and LANDAU et al. (2006) used the methodology 4 5 of BROSH et al. (1998) with HR and VO, measurements for 15-20 min throughout the day 6 7 and obtained results similar to those found in the 8 literature. However, according to PUCHALA et al. 9 (2007), the HR and energy expenditure of goats 10 consuming different quality diets varied within 24 11 h, thereby corroborating our observations.

12 The measurement time (up to 12 hours) 13 affected VO<sub>2</sub> (L/day) and VO<sub>2</sub> (mL/min/BW<sup>0.75</sup>). 14 From 15 h onwards, the parameters were similar to those obtained after 24 h (Table 4). Oxygen 15 16 consumption may have varied at the beginning of 17 the gas collection phase owing to the start of feeding and the initial stress associated with the chamber, 18 19 which resulted in increased  $O_2$  consumption. 20 VAN MILGEN et al. (1997) observed that 21 oxygen consumption varied according to animal 22 behavior when assessing O<sub>2</sub> consumption and CO<sub>2</sub> 23 production during the resting state, feeding and 24 physical activity in pigs. BARKAI et al. (2002) 25 and LANDAU et al. (2006) estimated oxygen consumption for 15-20 min at different times of the 26 27 day using the methodology of BROSH et al. (1998) 28 and observed no variation in oxygen consumption. 29 However, based on our observations, more accurate 30 measurements of the oxygen consumption require 31 longer measurement times.

32 Although it was possible to measure O<sub>2</sub>P 33 (mL/beat) for 9 h, the effect of time on O<sub>2</sub>P (mL/beat/ BW<sup>0.75</sup>) and HP<sub>O2P</sub> (Kcal/day and Kcal/day/BW<sup>0.75</sup>) 34 over the entire measurement period, demonstrated 35 that these parameters should be measured for 24 36 37 h when using the O<sub>2</sub>P methodology (Table 4). 38 This may be associated with variations in HR or 39 processes involving digestion and diet quality. The 40 roughage:concentrate ratio (40:60) explains the 24 h variations in HP<sub>02P</sub> because the degradation of non-41 fibrous carbohydrates is fast, whereas the digestion of 42 43 fibrous carbohydrates occurs more slowly owing to 44 the long lag time. 45

46 CONCLUSION

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48 The O<sub>2</sub>P-HR method is highly correlated 49 with the respirometry chamber methodology for estimating heat production in sheep; however, O<sub>2</sub>P-50 51 HR should be measured for 24 h to ensure greater accuracy. Sunflower meal inclusion levels did not affect heat production in the animals.

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#### DECLARATION OF CONFLICT OF INTEREST

The authors declare no conflict of interest. The founding sponsors had no role in the design of the study; in the collection, analyses, or interpretation of data; in the writing of the manuscript, and in the decision to publish the results.

#### **AUTHORS' CONTRIBUTIONS**

Conceptualization: LCG and ASC. Data acquisition: SSS, ASC and LCG. Design of methodology and data analysis: LCG and ASC. LCG, SSS, ASC and FSM prepared the draft of the manuscript. All authors critically revised the manuscript and approved of the final version.

#### BIOETHICS AND BIOSSECURITY COMMITTEE APPROVAL

The experimental procedures involving animals were approved by the Ethics Committee on Animal Use of the Universidade Federal de Minas Gerais (UFMG) under protocol No. 189/15.

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