INTRODUCTION

Metabolic acidosis is an acid-base imbalance that occurs in several diseases of ruminants, including acute ruminal lactic acidosis (ARLA) (SNYDER & CREDILLE, 2017), diarrhea, acetonemia, renal failure, hypovolemia, endotoxemia, and septicemic shock (CONSTABLE et al., 2016). Correction of metabolic acidosis is achieved by the administration of solutions containing alkalizing substances such as bicarbonate, which is the preferred buffer since it has an immediate effect (CONSTABLE, 2003; SNYDER & CREDILLE, 2017). However, rapid administration or administration of large amount can lead to adverse effects, such as iatrogenic alkalosis, hypernatremia, hypokalemia, and decreased ionized calcium (MUIR, 2017). Blood gas analysis is fundamental for the correct calculation of the required
amount of bicarbonate to be administered; however, this feature is not available for many professionals, especially those who work in the field.

Alternatively, metabolizable bases can also be used as they exert an alkalizing effect due to the consumption of H+ ions during their metabolism (CONSTABLE, 2003). Studies have already demonstrated the alkalizing potential of propionate, acetate, and lactate in cattle (NAYLOR & FORSYTH, 1986; LEAL et al., 2007). In Brazil, only lactate is reported in commercial electrolytic solutions such as lactated Ringer’s solution (LRS). Due to a low lactate concentration (28 mEq/L), the LRS alkalizing potential is reduced or absent in healthy animals (COSENZA et al., 2013; JUNQUEIRA et al., 2015), and this solution is not effective in correcting metabolic acidosis in diarrheic calves (NAKAGAWA et al., 2009) and sheep with ARLA (COSENZA et al., 2015).

In search of a safer alternative for the treatment of metabolic acidosis, an electrolyte solution containing 84 mEq/L of lactate was developed, and its alkalizing effect was verified in healthy sheep (FLAIBAN, 2010), calves (JUNQUEIRA et al., 2015), and horses (PINTO et al., 2018). This solution has been shown to be effective in correcting metabolic acidosis in sheep with ARLA (FLAIBAN, 2010), calves with diarrhea (JUNQUEIRA, 2012), and horses with hyperchloremic acidosis (ROMÃO et al., 2017). Despite the promising results, the effects of this solution have not yet been tested in goats. This study investigated the alkalizing potential of an electrolytic solution containing 84 mEq/L of lactate in healthy goats.

MATERIALS AND METHODS

Solutions

The crystalloid solutions presented in table 1 were used based on the composition of the commercial LRS (L28) and its osmolarity and sodium content. The lactate concentration was measured in triplicate (L84), and the chloride concentration was reduced. The bicarbonate solutions (B28 and B84) were equimolar to the lactate solutions and were used as the standard of comparison.

The solutions were prepared with commercial sterilized water in 1000 mL bottles (Water for injection: Halexistar Pharmaceutical Industry, Brazil). The components used in the preparation were sodium chloride (NaCl), potassium chloride (KCl), calcium chloride (CaCl2), sodium bicarbonate (NaHCO3), and sodium lactate, which were all pure for analysis (Synth; Labsynth, Brazil). L-lactate concentrations were measured and corresponded to 98% of the total lactate present. The solutions were prepared shortly before administration by adding the components in appropriate amounts or volume for each type of composition and finally homogenizing them. To avoid any contamination, aseptic care was rigorous and constant throughout solution preparation.

Experimental procedures

Six healthy, non-pregnant and non-lactating female Saanen young goats (13.95 ± 2.42 months of age) were enrolled in the study. The goats were housed in collective stalls and fed with water and coast-cross grass hay (Cynodondactylon) ad libitum, commercial ration (Cocari Goats adult; Cocari, Brazil) (300 g per animal, divided into two portions per day), and mineral salt (Sal Rural Maringá; SRM, Brazil).

The goats weighed 32.5 ± 3.5 kg, and a cross-over design was implemented, in which all six goats receiving the four solutions studied one at a time, with an interval of 4–5 days between infusions. Solutions assignments were randomly established. The solutions were infused intravenously, and the left jugular vein was kept catheterized (BD Angiocath®).
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The administered volume corresponded to 10% of body weight and the infusion rate was 33.3 mL/kg/h, totaling 3 h of continuous administration.

Venous blood and urine samples were collected immediately before starting the infusion and after initiation at 1.5h (half volume), 3h (end of infusion), 4.5h, and 6h. An extra sample of blood was collected at 24h from the beginning of the infusion. Blood samples were collected by puncturing the right jugular vein. Vacuum tubes (Laborvacum; Labor Import, Brazil) containing sodium fluoride were used, and the fluoridated plasma was obtained by centrifugation (10,000 rpm for 5 minutes) performed within a maximum of 10 minutes. The plasma was preserved by freezing (-20°C) until analysis. For hemogasometry, 3mL plastic syringes containing approximately 0.08 mL (400 IU) of sodium heparin (Hemofol; Cristália, Brazil) were used, and the samples were processed immediately. Urine samples were either obtained by spontaneous urination or induction by brief manual obstruction of the nostrils and mouth.

Physical examinations (PUGH et al., 2020) were performed every 90 minutes, until 6 hours after starting the infusion and 24 hours later. The animals were monitored for the occurrence of behavioral changes or any other manifestation during the experimental period.

Laboratory procedures

The blood pH, partial pressure of carbon dioxide (pCO₂), bicarbonate (HCO₃⁻), base excess (BE), sodium (Na⁺), chloride (Cl⁻), and potassium (K⁺) were measured using a gas analyzer (Omni C; Roche, Switzerland). Total plasma protein (TPP) concentrations were measured by a refractometer (PG1800; Gehaka, Brazil) and density by refractometry. In the frozen plasma, creatinine concentrations were determined by the enzymatic method (Dimension; Siemens). In the fresh urine samples, pH was determined using an electronic potentiometer (PG1800; Gehaka, Brazil) and density by refractometry. In the frozen samples, the creatinine concentrations were determined by the enzymatic method (Dimension; Siemens), L-lactate by colorimetry (Dimension; Siemens), and Na⁺, K⁺, and Cl⁻ by the selective ion electrode method (Dimension; Siemens).

The following variables were calculated using the corresponding formulas:

- Anion Gap (AG) (CONSTABLE, 1999): AG = (Na⁺ + K⁺) - (Cl⁻ + HCO₃⁻)
- Total concentration of non-volatile weak acids (A_wt) (CONSTABLE, 2002): A_wt = TPP (g/dL) × 3.6
- Percent change in plasma volume (%PV) (CARLSON & BRUSS, 2008): %PV = [(TPP₁ / TPP₂) - 1] × 100
- Urinary fractional excretions of electrolytes and L-lactate (GARRY et al., 1990): FE_a = (urinary concentration of a × plasma creatinine / plasma concentration of a) × 100, where a is the substance excreted.

Statistical methods

A bifactorial variance analysis was conducted to test the effects of time (during and after the infusion of solutions) and the type of solution administered, and to verify the existence of interaction between the two factors (time and type of solution). When the F statistic was significant, Tukey’s test was used. A probability of error of 5% was allowed. SigmaStat for Windows 3.1 program was used for statistical analysis.

RESULTS AND DISCUSSION

The L28 solution, equivalent to LRS, did not affect the pH, pCO₂, K⁺, Cl⁻, and SID values (P > 0.05) but caused slight elevations of HCO₃⁻ (P < 0.001), BE (P < 0.001), and Na⁺ (P < 0.001) at the end of the infusion (Figures 1 and 2), which returned to the original values at 4.5h. The L28 solution did not interfere with the electrolyte and acid-base balance of healthy goats when administered at a volume corresponding to 10% of body weight. Similar results were observed in healthy sheep, calves, and horses (COSENZA et al., 2013, JUNQUEIRA et al., 2015). These results contradicted the concept that LRS has an alkalizing capacity (CONSTABLE, 2003; JONES & NAVARRE, 2014) and that it could be used in the treatment of metabolic acidosis in calves (SMITH & BERCHTOLD, 2014). LRS was not able to correct the metabolic acidosis of calves with diarrhea (NAKAGAWA et al., 2009) and sheep with ALRA (COSENZA et al., 2015). As LRS has a plasma-like electrolyte composition, it is expected that it will not cause electrolyte and iatrogenic acid-base changes (COSENZA et al., 2013; JUNQUEIRA et al., 2015), which was also observed in this study. The alkalizing capacity of LRS is reduced due to the low effective SID₃, which has the same concentration as that of lactate (28 mEq/L) and does not remain in the plasma.
as a strong anion after its metabolism (CONSTABLE, 2014; MUIR, 2017).

The L84 solution, conversely, produced intense elevations of pH, HCO₃⁻ and BE ($P < 0.001$), starting from the middle of the infusion to reaching its peak toward the end. The pCO₂ remained unchanged ($P > 0.05$) (Figure 1). With the L84 infusion, Na⁺ concentration increased ($P < 0.001$), while that of K⁺ ($P < 0.001$) and Cl⁻ ($P = 0.02$) decreased, causing elevation of SID₃ ($P = 0.003$) and hypokalemia, which extended from the middle of the infusion to 1.5 h after its end (4.5 h) (Figure 2). These variations were transient, with return to the original values within 6 h for Na⁺, Cl⁻, and SID₃ or the next day (24 h) for the other variables.

Acid-base and electrolyte changes caused by solutions L28 and L84 were equivalent ($P > 0.05$) to those produced by solutions B28 and B84, respectively. The only exception was in chloremia and SID₃, in which the variation obtained from L84 infusion was different from that obtained from B84 infusion. Naturally, the Cl⁻ concentrations were lower and the SID₃ concentrations were higher after the end of the L84 infusion since it contained less Cl⁻ than that of the B84 solution.

At the end of L84 infusion, elevations of HCO₃⁻ and BE values were accentuated, and the blood pH reached the upper physiological limit, admitted as 7.54 (CARLSON & BRUSS, 2008). The compensatory pulmonary response to hypercapnia has not been well characterized in goats. The type and magnitude of the iatrogenic acid-base imbalance were similar to those observed previously in healthy sheep (FLAIBAN, 2010) and calves (JUNQUEIRA et al., 2015). An absence of compensatory hypercapnia was observed in sheep, contrary to that observed in calves. As already FLAIBAN (2010) speculated, it might be possible that, in goats, renal excretion mechanisms promote the rapid correction of alkalosis and make respiratory compensation minimally necessary. In fact, iatrogenic alkalosis was completely corrected on the day after the infusion (24 h), but at 3 h after the end of infusion (6 h), it was already attenuated.

The increase in Na⁺ concentration and decrease in Cl⁻ concentration observed at the end of L84 infusion occurred because of its composition, whose Cl⁻ concentration was much lower than that of plasma. Conversely, hypopotassemia, observed
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The plasma L-lactate concentration decreased with the infusion of solutions L28, B28, and B84 \((P < 0.001)\). The decrease in the plasma L-lactate concentration in animals of the L28 group occurred due to rapid lactate metabolism and mainly due to the hemodilution effect observed after infusion. Conversely, administration of L84 solution resulted in an elevation of L-lactate concentration during the infusion \((P < 0.001)\), which reverted to baseline value 1 h and 30 minutes after its end (4.5 h) (Figure 3). In the middle of the B28 and B84 infusions, L-lactate concentrations were found to be lower than those of the L28 and L84 solutions \((P < 0.001)\). L-lactate FE was slightly elevated \((P < 0.001)\) after the end of the infusion (4.5 and 6 h), regardless of the type of solution administered \((P = 0.525)\). The slight increase in the L-lactate FE after L28 and L84 solutions infusions can be explained by the renal excretion of part of the L-lactate, which was not metabolized. However, the reason for the increase in L-lactate FE after the infusion of solutions containing sodium bicarbonate (B28 and B84) is not known. This result is different from that observed in healthy sheep and calves, which showed no difference in L-lactate FE before, during, or after the solution infusions (FLAIBAN, 2010; JUNQUEIRA et al., 2015). Healthy horses, when receiving L84 solution at a speed of 16.66 mL/kg/h, showed an increase in L-lactate FE at the end of the infusion, with a return to baseline values 6 h after the end of the infusion (PINTO et al., 2018).

Infusion of solutions at a volume corresponding to 10% of body weight resulted in a reduction in TPP values \((P < 0.001)\), A\(_{Na+}\) \((P < 0.001)\), and AG \((P = 0.008)\), and an increase in %PV \((P < 0.001)\) (Figure 3). These effects occurred without distinction between the

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solutions ($P > 0.05$) and were transient, observed from the middle of the infusion, and returned to the original values 3 h after the end (6 h). The observed reductions in TPP, $A_{\text{tot}}$, and AG occurred due to hemodilution caused by the administration of the solutions in a volume corresponding to 10% of body weight in healthy, non-dehydrated animals. The elevation in % PV confirmed this. According to the strong ion model, $A_{\text{tot}}$ represents the negative charge of plasma proteins and has an influence on the acid-base balance (CONSTABLE, 2014; MUIR, 2017). Hypoproteinemia, reflected in the reduction of this variable, is one of the causes of metabolic alkalosis (CONSTABLE, 2014). However, the decrease in $A_{\text{tot}}$ should not be considered as an additional cause of the alkalizing effect observed in goats, since it also occurred after the administration of all the solutions used in the present study.

The frequency of urination increased during the infusion. Urine presented a lower density ($P < 0.001$) from the middle of the infusion, and the original values were recovered 3 h after the end of infusion (6 h). The pH of the urine did not change over time ($P=0.059$); however, at 4.5 h, the L84 solution caused higher values than those of L28 ($P=0.014$).
An increase in urinary pH would be expected due to the elimination of bicarbonate owing to the renal mechanism for correcting the acid-base imbalance caused by infusing alkaline solutions. Urine from adult ruminants is naturally alkaline, and additional alkalization is not always apparent after the administration of alkalizing solutions (CARLSON & BRUSS, 2008). Similar results have been reported for healthy sheep (FLAIBAN, 2010). Na⁺ and Cl⁻ FE increased \((P < 0.001)\) after the end of L84 infusion, and K⁺ FE decreased during the infusion \((P < 0.001)\) of all solutions studied (Figure 4). Except for pH, the variables studied in urine were not affected by the type of solution administered \((P > 0.05)\). All iatrogenic electrolyte changes were transient and corrected by the kidneys through selective excretion of ions. An increase in FE of Na⁺ and Cl⁻ is necessary to maintain homeostasis, minimizing the imbalance caused by the administration of excess electrolytes (LUNN & MCGUIR, 1990). The decrease in K⁺ FE during infusion, with return to original values at 4.5 h, occurred in response to hemodilution, hyperhydration, and the need to excrete excess liquid in the studied animals. This result is different from that observed in studies in sheep (FLAIBAN, 2010) and horses (PINTO et al., 2018), which showed elevated K⁺ excretion, and in calves (JUNQUEIRA et al., 2015), which did not show any variation.

During the period of study, goats remained alert and had appetite. No apparent clinical side effects were observed during or after infusions of the solutions. The values of body temperature, heart rate, respiratory rate, and frequency of ruminal movements did not change during or after the infusions \((P > 0.05)\) or due to the type of solution infused \((P > 0.05)\).

The alkalizing effects of L84 solution have previously been demonstrated in studies conducted in healthy sheep (FLAIBAN, 2010) and calves (JUNQUEIRA et al., 2015). In these studies, L84 solution was also administered in a volume corresponding to 10% of body weight, but the infusion rate was lower (20 and 25 mL/kg/h). Another distinction from the previous studies is that, unlike sheep and calves that were followed for only 2 and 2.5 h after the end of the infusion, the goats were followed for a longer period of time, which allowed us to verify the time in which the iatrogenic imbalance was reversed. Another study, using the L84 solution...
in healthy horses, with different infusion rates (16.66 or 8.33 mL/kg/h), proved the alkalinizing capacity of the solution in this species; however, it was revealed that its magnitude is dependent on the infusion rate (PINTO et al., 2018).

According to the simplified strong ion model (CONSTABLE, 1999; MUIR, 2017), plasma electrolytes influence acid-base balance and blood pH, and plasma SID₃ is the representative variable for this event. An increase in the alkalinizing capacity of L84 solution could be achieved by tripling the concentration of lactate and decreasing the concentration of CI. Thus, L84 administration resulted in a reduction in chloremia, which increased SID₃ and consequently generated iatrogenic metabolic alkalosis. The increase in SID₃ at the end of L84 infusion was also demonstrated in healthy sheep, calves, and horses (FLAIBAN, 2010; JUNQUEIRA et al., 2015; PINTO et al., 2018). The effective SID₃ of the L84 solution is equivalent to its lactate concentration (84 mEq/L). With the metabolism of this anion, the impact of the reduced CI concentration assumes the greatest importance because CI becomes the only anion infused to exert influence on chloremia and plasma SID₃. Theretewith which iatrogenic hyperlactatemia was reversed reinforced previous observations in sheep and healthy calves that received L84 solution (FLAIBAN, 2010; JUNQUEIRA et al., 2015) and proved that, in healthy goats too, the metabolization of infused lactate is rapid. Therefore, reduction of chloremia, elevation of plasma SID₃, and alkalinizing effect are already established rapidly during L84 infusion.

The effective SID, total volume and rate of solution administration, and the metabolism of the anions present in the solution are the main determinants of the solution’s effect on blood pH (MUIR, 2017). The L84 solution was effective in correcting metabolic acidosis in sheep with ARLA (FLAIBAN, 2010), in calves with diarrhea (JUNQUEIRA, 2012) and in horses with hyperchloremic acidosis (ROMÃO et al., 2017), without causing any side effects. Therefore, this study indicated that the L84 electrolyte solution can be used in the treatment of metabolic acidosis, as long as the organism’s ability to metabolize infused lactate is not compromised.

CONCLUSION

The electrolytic solution containing 84 mEq/L of lactate produced iatrogenic alkalinization when infused into healthy goats, without causing side effects. Future studies should be conducted to clarify the safety and efficacy of the solution in sick goats.

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BIOETICS AND BIOSSECURITY COMMITTEE APPROVAL

The project was approved by the Committee on Ethics in the Use of Animals of the Universidade Estadual de Londrina, under the protocol CEUA/UEL 132/2012.

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

PEREIRA, P.F.V performed the experiment, provided clinical data, carried out laboratory analysis and wrote the manuscript. ROMÃO, F.T.N.M.A, CURTI, J.M and CAMILO, S.L.O performed the experiment and carried out laboratory analysis. FLAIBAN, K.K.M.C carried out laboratory analysis and revised the manuscript. LISBÔA, J.A.N designed and supervised the experiment, performed statistical analyses of experimental data and revised the manuscript.

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