Effects of *Lactobacillus buchneri* on the nutritive value of sugarcane silage for finishing beef bulls

Patrick Schmidt², Luiz Gustavo Nussio³, Oscar Cézar Müller Queiroz⁴, Mateus Castilho Santos³, Maity Zopollatto², Sérgio Gil de Toledo Filho³, João Luiz Pratti Daniel³

1 Funded by FAPESP, CAPES, and CNPq.
2 Departamento de Zootecnia, Universidade Federal do Paraná, Curitiba, PR, Brasil.
3 Departamento de Zootecnia, Escola Superior de Agricultura “Luiz de Queiroz”, Piracicaba, SP, Brasil.
4 Department of Animal Sciences, University of Florida, Gainesville, USA.

**ABSTRACT** - *Lactobacillus buchneri* 40788 and the fibrolytic enzymes β-glucanase and xylanase were applied to chopped sugarcane to study their effects on the nutritive value of silage. Sugarcane was mechanically harvested after 14 mo of growth and treated without (control) or with *L. buchneri* at a theoretical application rate of $5 \times 10^4$ cfu/g, $1 \times 10^5$ cfu/g, or $1 \times 10^6$ cfu/g plus enzymes. Forage was packed into farm-scale bag silos (40 t/silo) and stored for 92 d. Fifty-six bulls (32 Nellore and 24 Charolais × Nellore) were housed in 20 collective pens and fed diets comprising (dry matter [DM] basis) 458 g/kg sugarcane silage and 542 g/kg concentrates for an 84-d period. Treated silages had higher concentrations of acetic acid and lower concentrations of ethanol. Total mixed rations (TMR) containing inoculated silages exhibited significantly lower neutral detergent fiber (NDF) concentration and, consequently, higher *in vitro* DM digestibility (IVDMD). Thus, animals fed TMR containing treated silages spent less time chewing per day and per kilogram of DM intake (DMI), even at higher DMI levels. Nonetheless, the intake of NDF was similar across treatments (0.77 to 0.79 kg/100 kg BW) but markedly lower than the value reported for traditional forages. Average daily gain was significantly greater for animals fed TMR based on inoculated silages due to the higher DMI (14% on average) and the higher energy content of the diets, as indicated by the higher feed efficiency (12% on average). The dose of inoculants used and the addition of fibrolytic enzymes had no significant effects on silage parameters or animal performance. Therefore, inoculation of *L. buchneri* during sugarcane ensilage can alter the fermentation pattern by increasing acetic acid yield, reducing silage nutrient losses, and improving feed efficiency by bulls.

Key Words: feed efficiency, heterolactic bacteria, *Saccharum officinarum* L., voluntary feed intake

**Introduction**

Fresh chopped sugarcane is widely used for feeding beef and dairy cattle because its harvesting period coincides with the period of pasture shortage in Brazil. However, to avoid daily harvesting, chopping, and hauling and to prevent crop loss by accidental fire, this forage could be ensiled. In addition, considering that sugarcane is a semi-perennial tropical grass, its field lifespan may be prolonged by uniform harvesting and post-harvesting management.

Ensiling sugarcane results in the conversion of most of the water-soluble carbohydrates (WSC) into fermentation end-products, which are characterized by high levels of volatile organic compounds, mainly ethanol (Kung Jr. and Stanley, 1982; Pedroso et al., 2005). Although the gross energy is almost completely recovered during alcoholic fermentation, large amounts of DM and net energy are lost (McDonald et al., 1991; Daniel and Nussio, 2011). Furthermore, ethanol is metabolized to acetate in the rumen with concomitant methane formation (Yoshii et al., 2005), which has a negative impact on the environment. Due to the undesirable characteristics of the natural fermentation of sugarcane, additives have been recommended to inhibit epiphytic yeast populations and mitigate alcohol synthesis in sugarcane silages (Pedroso et al., 2008).

A heterofermentative lactic acid bacterium, *Lactobacillus buchneri*, has been studied as an inoculant to improve the preservation of sugarcane silage during both anaerobic storage and air exposure (Pedroso et al., 2008; Ávila et al., 2009). *L. buchneri* is known to produce acetic acid (Oude Elferink et al., 2001; Pahlow et al., 2003), which is a powerful antifungal agent (Danner et al., 2003) capable of decreasing ethanol production and improving the aerobic stability of silages (Ranjit et al., 2002; Reich and Kung Jr., 2010). Conversely, high concentrations of acetic acid in silages have the potential to reduce feed intake (Dinius et al., 1968; Hutchinson and Wilkins, 1971).
Several studies have reported increased DM recovery when *L. buchneri* was applied to sugarcane silages (Schmidt, 2009; Zopollatto et al., 2009). However, experiments involving animals fed sugarcane silage are scarce. The objectives of this study were to evaluate the effects of dose of *L. buchneri* (strain 40788) and its association with fibrolytic enzymes on the fermentation of sugarcane silage ensiled in farm-scale silos and on the performance of finishing beef bulls. We hypothesized that *L. buchneri* alone or associated with fibrolytic enzymes may improve the nutritive value of sugarcane silages for finishing beef bulls.

**Material and Methods**

Sugarcane variety RB 85-5536 was mechanically harvested (Colhiflex Mentamit®, Cajurú, Brazil) from one field after 14 mo of growth (first cut) in 2002-2003 crop year, to a theoretical cut of 8 mm, and packed into 4 bag silos (2.7 m i.d., Pacifil, Estância Velha, Brazil). At harvest, the content of soluble solids in the sugarcane juice was 21.8°brix, indicating that the crop was mature. The mean chemical composition (DM basis) of fresh sugarcane was 332 g/kg DM, 23 g/kg ash, 502 g/kg NDF, 290 g/kg ADF, and 41 g/kg CP, and in vitro DM digestibility (IVDMD) was 615 g/kg.

The forage placed in one bag was sprayed with water (Control), whereas the remaining three bags were treated as follows: (LLB) a low dose of *L. buchneri* 40788 (final application rate of 5 × 10^4 cfu/g of fresh forage); (HLB) a high dose of *L. buchneri* 40788 (final application rate of 1 × 10^5 cfu/g of fresh forage); and (HLBE) a high dose of *L. buchneri* 40788 plus fibrolytic enzymes (final application rate of 1 × 10^5 cfu/g plus β-glucanase at 32,340 IU/t and xylanase at 40,165 IU/t of fresh forage; Lallemand Animal Nutrition, Milwaukee, WI). Aqueous solutions of the additives were sprayed onto the forages at a rate of 2.2 L/t. Approximately 40 t were packed into each bag.

After 92 d of storage, the bags were opened and the silages were used to prepare four diets for feeding beef bulls. The diets comprised (DM basis) 458 g/kg sugarcane silage, 314 g/kg dried citrus pulp, 203 g/kg corn gluten feed, 14 g/kg urea, and 11 g/kg mineral and vitamin mix. The mineral and vitamin mix contained 40 g/kg Na; 65 g/kg S; 9 ppm Co; 1,000 ppm Cu; 600 ppm Mn; 2,500 ppm Zn; 50 ppm I; 10 ppm Se; 350,000 IU/kg vitamin A; 30,000 IU/kg vitamin D; 1,800 IU/kg vitamin E; and 25 g/kg monensin. Dietary requirements of Ca and P were met with concentrate ingredients (NRC, 1996). The four diet treatments were named in the same way as the silages described above.

Fifty-six bulls (32 Nellore and 24 Charolais × Nellore, 15 to 18 mo old) from the University of São Paulo herd were stratified by breed and BW and randomly housed in 20 covered pens (21 m², concrete floor); two or three animals were housed in each pen. Fresh water was available at all times, and the animals were cared for using accepted protocols (FASS, 2010). The bulls were de-wormed with ivermectin (200 µg/kg BW; Ivomec Merial®, Paulínia, Brazil) and acclimated to the facilities for 21 d. During the adaptation period, the bulls were fed a diet containing 542 g/kg concentrates (as above) and 458 g/kg sugarcane silages (1/4 from each treatment, DM basis). Body weight was recorded after 12 h of fasting at the beginning and end of the study (84-d period). Initial BW was 426±54 kg (mean ± SD) for Nellore and 513±43 kg for Charolais × Nellore bulls.

Silages were feed-out, mixed with concentrates, and animals were fed TMR once a day targeting 100 g/kg of orts in the next day. Feed intake was determined by the difference between the amount of feed offered and refused every day. The DMI, average daily gain (ADG), and feed efficiency (ADG:DMI) were estimated for the 84-d feeding period. Digestive behavior was recorded on day 48 of the experiment by visual observation of the animals throughout a 24-h period. Eating and ruminating activities were recorded at 10-min intervals and a 24-h pattern was estimated considering a constant behavior between observations (Maekawa et al., 2002). Chewing (eating + ruminating) per kilogram of DM and NDF was calculated using measurements of DMI and NDF intake recorded on the same day.

Silage, TMR, and orts were sampled weekly (n = 12) and dried in an air-forced oven at 55 °C for 72 h. Aliquots of silages were also used to determine the particle size (Lammers et al., 1996) and prepare aqueous extracts (Kung Jr. et al., 1984).

Dried samples were ground through a 1-mm screen (Wiley mill), and sub-samples were analyzed for the following: DM in an air-forced oven at 105 °C for 24 h (AOAC, 1980); CP by the Dumas method (Wiles et al., 1998); ash (AOAC, 1980); and NDF and ADF (sequential and expressed inclusive of residual ash; Van Soest et al., 1991). Hemicellulose was calculated as NDF minus ADF. The IVDMD was determined using the Tilley and Terry method as modified by Goering and Van Soest (1970).

Aqueous extracts from silages were analyzed for pH, VFA (Palmquist and Conrad, 1971), lactic acid (Pryce, 1969), and WSC (Dubois et al., 1956). Ethanol concentration was determined using a biochemistry analyzer (YSI 2700 Biochemistry Analyzer, Yellow Springs, OH) that uses
membrane-immobilized ethanol oxidase (EC 1.1.3.13) (Taylor et al., 1999).

Statistical analysis was performed by the MIXED procedure of SAS (Statistical Analysis System, version 9.2) using the following model: $y_{ij} = \mu + \alpha_i + \beta_j + \epsilon_{ij}$, in which $\mu$ = overall mean; $\alpha_i$ = random effect of block ($i = 1$ to $5$); $\beta_j$ = fixed effect of treatment ($j = 1$ to $4$); and $\epsilon_{ij}$ = residual error. The block effect included the variations of animal breed and initial BW. Pens were considered as the experimental units. Degrees of freedom of treatment were partitioned into three single degree of freedom orthogonal contrasts: additive application effect (Control vs. LLB+HLB+HLBE), additive dose effect (LLB vs. HLB+HLB+HLBE), and enzyme application effect (HLB vs. HLBE). Contrasts were declared significant at $P \leq 0.05$.

**Results and Discussion**

There were no great alterations in chemical entities among silages (Table 1), although inoculated silages had higher concentrations of acetic acid and lower concentrations of ethanol. *L. buchneri* is known to produce acetic acid (Oude Elferink et al., 2001), which is a powerful antifungal agent (Danner et al., 2003) capable of inhibiting yeast and reducing ethanol formation and DM loss during silage fermentation (Ranjit et al., 2002; Pedroso et al., 2008; Ávila et al., 2009). In the present study, *L. buchneri* increased the acetic acid content by 31.5% and decreased the ethanol concentration by 70.8%. Nevertheless, ethanol concentrations were much lower than those reported in the literature (Zopollatto et al., 2009; Daniel et al., 2013), probably because of partial volatilization during silage feedout.

Unexpectedly, the DM contents of silages were equivalent to that of fresh sugarcane before ensiling. The conversion of WSC to fermentation end-products generates gases and water (McDonald et al., 1991), which ultimately increase the moisture content. Ensiling sugarcane under laboratory conditions normally decreases DM concentrations (Schmidt, 2009). In farm-scale silos, however, this effect is not always evident if the losses of DM and moisture occur at similar magnitudes (Pedroso et al., 2006). Furthermore, most of the volatile compounds may have been lost during the oven-drying of the silage samples in the laboratory (McDonald and Dewar, 1960; Daniel and Nussio, 2011), thus leading to underestimation of the DM content and reinforcing the extensive occurrence of water loss from the silo working face.

Inoculant dose and addition of fibrolytic enzymes led to silages with similar composition. Several authors have reported seeing no effect of fibrolytic enzymes on tropical silages (Loures et al., 2005; Avellaneda-Cevallos et al., 2009). Sugarcane crop typically contains a high content of WSC. Therefore, the small amount of sugars provided by cell wall hydrolysis would not be expected to change sugar abundance and, thus, the fermentation process.

Rations containing inoculated silages exhibited (numerically) lower NDF and, consequently, higher IVDMD (Table 2). Similarly, the higher DM content may indicate that nutrients are better preserved in treated silages (Schmidt, 2009). These alterations suggest the higher nutritive value of sugarcane inoculated with additives as indicated by animal performance.

Ingestive behavior was recorded to verify the possible adverse effects of silage inoculation on the eating pattern.
because high levels of acetic acid have been associated with decreased intake (Dinius et al., 1968). In the present study, silage inoculation with L. buchneri did not significantly affect eating (P = 0.113) or rumination (P = 0.233; Table 3). However, because of the higher content of NDF, animals fed TMR that contained control silage spent more time chewing per day (P = 0.021) and per kilogram of DMI (P<0.001), even with a lower DMI (see below). A dose effect was observed for rumination (P = 0.011), but the reason for this was unclear. As anticipated, chewing/kg NDF intake was unchanged by treatments (P = 0.177). Therefore, enzyme application was unable to alter the ability of fiber to stimulate chewing (P = 0.916).

Bulls fed TMR containing sugarcane silage inoculated with L. buchneri had higher DMI (P = 0.006; Table 4). Several studies have shown that when cattle or sheep are fed silages treated with L. buchneri, DMI is not affected (Driehuis et al., 1999; Taylor et al., 2002; Ranjit et al., 2002; Kung et al., 2003). However, in the present study, the inoculants were effective in mitigating NDF accumulation caused by sugar oxidation during the ensiling process and improved the DMI. Indeed, the control treatment limited DMI with rumen fill from dietary fiber. Nonetheless, the NDF intake was similar across treatments (0.77 to 1.2 kg/100 kg BW; P = 0.555) but substantially lower than the value reported for traditional forages (1.0 to 1.2 kg/100 kg BW; Mertens, 1994; Krizsan et al., 2010). Feed intake constraint was consistent with the higher chewing activity associated with sugarcane-based diets compared with conventional roughage (Corrêa et al., 2003; Costa et al., 2005).

The average daily gain (ADG) was significantly greater for animals fed TMR based on inoculated silages (P = 0.003) (Table 4) due to the higher DMI (14% on average) and the higher energy content of the diets, as indicated by the superior feed efficiency (12% on average). In addition to increased ADG, animals fed TMR that contained inoculated silages exhibited higher final BW and total BW gain across the 84-d finishing period. The application rate of the inoculants and the addition of fibrolytic enzymes did not affect animal performance (P>0.119). An earlier study carried out by Pedroso et al. (2006) reported an increase of 31.9% in the ADG of Holstein heifers fed a TMR that contained sugarcane inoculated with L. buchneri 40788 (1.24 kg/d) compared with animals fed a TMR that was based on untreated silage (0.94 kg/d). In the current trial, silage DM was determined by oven drying; thus, the DM content may have been underestimated, leading to underestimation of DMI and overestimation of feed efficiency (McDonald and Dewar, 1960; Daniel and Nussio, 2011).

It is important to emphasize that the dietary level of forage adopted in the current study was within the practical range under Brazilian conditions (Milhen et al., 2009) but is much greater than the concentration of roughage used in feedlots in the U.S. (Vasconcelos and Galyean, 2007). Therefore, one can expect less improvement in animal performance resulting from the use of silage additives when TMR containing less forage are fed.

Table 3 - Influence of L. buchneri inoculation of sugarcane silages on the ingestive behavior of beef bulls

<table>
<thead>
<tr>
<th>Item</th>
<th>Control</th>
<th>LLB</th>
<th>HLB</th>
<th>HLBE</th>
<th>SEM</th>
<th>P-contrasts&lt;sup&gt;1&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>A</td>
</tr>
<tr>
<td>Eating (min/d)</td>
<td>233</td>
<td>218</td>
<td>191</td>
<td>212</td>
<td>15.0</td>
<td>0.113</td>
</tr>
<tr>
<td>Ruminating (min/d)</td>
<td>546</td>
<td>510</td>
<td>552</td>
<td>544</td>
<td>9.7</td>
<td>0.233</td>
</tr>
<tr>
<td>Chewing (min/d)</td>
<td>785</td>
<td>737</td>
<td>733</td>
<td>747</td>
<td>15.9</td>
<td>0.021</td>
</tr>
<tr>
<td>Chewing/DMI (min/kg)</td>
<td>107</td>
<td>80</td>
<td>80</td>
<td>85</td>
<td>4.50</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Chewing/NDF intake (min/kg)</td>
<td>201</td>
<td>171</td>
<td>184</td>
<td>183</td>
<td>9.87</td>
<td>0.177</td>
</tr>
</tbody>
</table>

SEM - standard error of the mean; DMI - dry matter intake; NDF - neutral detergent fiber.

<sup>1</sup> Probability for contrasts. Additive effect: A = Control vs. (LLB+HLB+HLBE); Dose effect: D = LLB vs. (HLB+HLBE); Enzyme effect: E = HLB vs. HLBE.

Table 4 - Effect of L. buchneri inoculation of sugarcane silages on intake and growth performance of finishing beef bulls for 84 days

<table>
<thead>
<tr>
<th>Item</th>
<th>Control</th>
<th>LLB</th>
<th>HLB</th>
<th>HLBE</th>
<th>SEM&lt;sup&gt;1&lt;/sup&gt;</th>
<th>P-contrasts&lt;sup&gt;2&lt;/sup&gt;</th>
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<tr>
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<td></td>
<td></td>
<td></td>
<td></td>
<td>A</td>
</tr>
<tr>
<td>Initial body weight (kg)</td>
<td>462</td>
<td>466</td>
<td>466</td>
<td>467</td>
<td>29.7</td>
<td>0.576</td>
</tr>
<tr>
<td>Final body weight (kg)</td>
<td>531</td>
<td>562</td>
<td>548</td>
<td>552</td>
<td>26.4</td>
<td>0.027</td>
</tr>
<tr>
<td>Total body weight gain (kg)</td>
<td>68</td>
<td>87</td>
<td>81</td>
<td>85</td>
<td>4.7</td>
<td>0.003</td>
</tr>
<tr>
<td>Dry matter intake (kg/d)</td>
<td>7.69</td>
<td>8.77</td>
<td>8.87</td>
<td>8.59</td>
<td>0.41</td>
<td>0.006</td>
</tr>
<tr>
<td>Average daily gain (kg/d)</td>
<td>0.817</td>
<td>1.040</td>
<td>0.969</td>
<td>1.007</td>
<td>0.056</td>
<td>0.003</td>
</tr>
<tr>
<td>ADG:DMI</td>
<td>0.104</td>
<td>0.120</td>
<td>0.110</td>
<td>0.118</td>
<td>0.009</td>
<td>0.028</td>
</tr>
</tbody>
</table>

SEM - standard error of the mean; ADG - average daily gain; DMI - dry matter intake.

<sup>1</sup> No. of pens = 20.

<sup>2</sup> Probability for contrasts. Additive effect: A = Control vs. (LLB+HLB+HLBE); Dose effect: D = LLB vs. (HLB+HLBE); Enzyme effect: E = HLB vs. HLBE.
Conclusions

Inoculating sugarcane with *L. buchneri* 40788 at ensiling can alter the fermentation process by increasing acetic acid production. Due to the antifungal properties of acetic acid, the total mixed ration containing treated silages have higher nutritive value. Bulls fed a total mixed ration containing sugarcane silages inoculated with *L. buchneri* 40788 eat greater amounts of dry matter and gain more body weight than bulls feeding untreated silage. Furthermore, silages treated with *L. buchneri* 40788 improve feed efficiency. The dose of *L. buchneri* and the addition of fibrolytic enzymes have no significant effects on silage parameters or animal performance.

References


