#### Note

# Production of ethanol from mesquite [Prosopis juliflora (SW) D.C.] pods mash by Zymomonas mobilis in submerged fermentation

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ABSTRACT: Mesquite [*Prosopis juliflora* (SW) D.C.], a perennial tropical plant commonly found in Brazilian semi-arid region, is a viable raw material for fermentative processes because of its low cost and production of pods with high content of hydrolysable sugars which generate many compounds, including ethanol. This study aimed to evaluate the use of mesquite pods as substrate for ethanol production by *Z. mobilis* UFPEDA-205 in a submerged fermentation. The fermentation was assessed for rate of substrate yield to ethanol, rate of ethanol production and efficiency of fermentation. The very close theoretical (170 g L<sup>-1</sup>) and experimental (165 g L<sup>-1</sup>) maximum ethanol yields were achieved at 36 h of fermentation. The highest counts of *Z. mobilis* UFEPEDA-205 (both close to 6 Log cfu mL<sup>-1</sup>) were also noted at 36 h. Highest rates of substrate yield to ethanol (0.44 g ethanol g glucose<sup>-1</sup>), of ethanol production (4.69 g L<sup>-1</sup> h<sup>-1</sup>) and of efficiency of fermentation (86.81%) were found after 30 h. These findings suggest mesquite pods as an interesting substrate for ethanol production using submerged fermentation by *Z. mobilis*.

Key words: renewable sources, bioconversion, fermentative parameters

# Produção de etanol a partir do mosto de vagens de algaroba [Prosopis juliflora (SW) D.C.] por Zymomonas mobilis em fermentação submersa

RESUMO: A algaroba [*Prosopis juliflora* (SW) D.C.] é uma planta tropical perene comumente encontrada no semi-árido brasileiro e apresenta-se como matéria-prima viável para o processo fermentativo por possuir baixo custo e para produzir vagens que contém um elevado teor de açúcares hidrolisáveis, os quais podem gerar diversos compostos, incluindo etanol. Avaliou-se o uso de vagens de algaroba como substrato para produção de etanol por *Z. mobilis* UFPEDA-205 por meio de fermentação submersa. O processo fermentativo foi avaliado por meio da mensuração da taxa de conversão de substrato em etanol, taxa de produção de etanol e eficiência de fermentação. Os valores muito próximos encontrados para o fornecimento máximo teórico (170 g L<sup>-1</sup>) e experimental (165 g L<sup>-1</sup>) de etanol foram alcançados após 36 h de fermentação. O valor de contagem experimental de *Z. mobilis* UFPEDA-205 (próximo a 6 Log cfu mL<sup>-1</sup>) foi encontrado após 36 h de fermentação. As mais elevadas taxas de conversão de substrato para etanol (0,44 g ethanol g glucose<sup>-1</sup>), de produção de etanol (4,69 g L<sup>-1</sup> h<sup>-1</sup>) e de eficiência de fermentação (86,81%) foram encontrados depois de 30 h. Conclui-se que as vagens de algaroba apresentam potencial como substrato emergente para produção de etanol por *Z. mobilis* por meio de fermentação submersa.

Palavras-chave: fontes renováveis, bioconversão, parâmetros fermentativos

### Introduction

Fuels obtained from renewable resources have deserved a great deal of interest during the past decades mainly due to concerns about fossil fuels depletion. Research efforts have been multiplied in the last years as a consequence of constant increasing costs and environmental impact derived from the use of crude-based fuels (Gray et al., 2006; Prasad et al., 2007). Zymomonas mobilis is a Gram-negative, facultative anaerobic that ferments glucose, fructose, and sucrose as carbon sources (Viikari, 1998). These carbohydrates are metabolized via the same biochemical route, the Entner-Doudoroff pathway (Paula et al., 2007). Z. mobilis is a promising alternative to yeast in the synthesis of ethanol. In comparison with yeast, Z. mobilis has a higher tolerance to ethanol and better kinetic characteristics such as higher specific substrate uptake, higher ethanol synthesis rate and higher substrate yield to ethanol. Moreover, it has advantages for the fermentation of glucose to ethanol that include a high yield of ethanol from consumed glucose and a high specific rate of ethanol production (Joachimsthal et al., 1998; Shene and Bravo, 2001; Tano and Buzato, 2003). The metabolic activity of *Z. mobilis* depends on the strain and carbon source, while many by-products can be produced during the fermentation of sucrose, such as phenol, lactic acid, higher alcohols, acetaldehyde, methanol and levan (Kalneniekis et al., 2000; Borsari et al., 2006).

Prosopis juliflora (SW) D.C., Leguminosae, a popular plant known as mesquite, is native to Central and South America and has spread to North America. Mesquite has great potential for use as a multipurpose tree in different parts of the world in comparison to several native and exotic species (Kailappan et al., 2000; Deans et al., 2003). Mesquite pods present a high amount of carbohydrates (Batista et al., 2002). Pod production per tree can vary from a few kg to over 400 kg and is highly dependent on moisture availability to the plant (Riveros, 1992). In the northeast region of Brazil, mesquite trees cover 150.000 ha (Tabosa et al., 2000).

The aim of this study was to assess the use of mesquite pods as substrate for ethanol production by Z. *mobilis* UFPEDA-205 in a submerged fermentation.

#### Material and Methods

Strain of Z. mobilis UFEPEDA-205 used in this study was supplied by the Department of Antibiotics, Federal University of Pernambuco, Recife, Brazil. Stock cultures were kept in slopes Standard Swings and De Ley – SSDL agar (glucose 20.0; yeast extract 5.0; agar 15 g L<sup>-1</sup>) (Swings and De Ley, 1977) under refrigeration. For experimental assays, Z. mobilis were grown in 50 mL of SSDL broth at 37°C. After 48 h incubation, 5 mL of the culture was added to flasks containing 95 mL of the same growth media and allowed to grow at room temperature for 24 h under rotation (150 rpm).

Liquefied mash was prepared using healthy (with no infection sign) mesquite pods. Mesquite pods used in this study presented moisture 5.8; total sugars 56.5; reducing sugars 4.6; total fiber 7.2; proteins 9.0; fat 2.1; and ashes  $0.2 \text{ g} 100 \text{ g}^{-1}$  (Silva et al., 2007). Pods were dried at 45°C for 18 h, followed for grounding in hammer mill with a #4 screen to get the appropriate grind size. Hydrated mash was prepared at 30 g 100 g<sup>-1</sup> of distilled water. To prepare the mash, ground mesquite was slowly added to distilled water in a constant agitation. After the addition of the proper ground amount, the mash was heated to 50°C, maintained at this temperature for 1 h and submitted to centrifugation (3000 rpm for 15 minutes). The supernatant was vacuumed filtered using Whatman n. 1 and autoclaved at 121°C for 15 min. After that, the mash was cooled to room temperature and aliquots were aseptically dispensed in sterile Erlenmeyer flasks for fermentation.

The mash used for fermentations presented total sugars (sucrose) 16.1; reducing sugars 3.99; total fiber 3.99; proteins 2.16; fat 0.63; tannins 0.09; and ashes 0.2 g  $100 \text{ g}^{-1}$ . It was reported a total soluble solids value of  $18^{\circ}$ Brix (Silva et al., 2007).

Submerged fermentation of mesquite hydrated mash by Z. mobilis UFEPEDA-205 was analyzed. A 100-mL aliquot of mesquite mash (added with 10 g L<sup>-1</sup> (NH<sub>4</sub>)<sub>2</sub>SO<sub>4</sub> and 2 g L<sup>-1</sup> KH<sub>2</sub>PO<sub>4</sub>) was aseptically distributed in sterile 250-mL Erlenmeyer flasks and inoculated with a 24 h-old culture (approximately 10<sup>8</sup> cfu mL<sup>-1</sup>). The flasks were incubated at room temperature (28 ± 1°C) under static condition. During 72 h of fermentation, the mash was analyzed for pH, glucose concentration, bacterial count and ethanol concentration. The fermentation was carried out in triplicate and the results were expressed as average of the parallel assays.

Kit Glicose PAP – Liquiform (Labtest Diagnóstica, Minas Gerais, Brazil) was used to measure the glucose concentration (g L<sup>-1</sup>), while the pH value was found using a Micronal B474 digital pHmeter. The growth of Z. *mobilis* was evaluated by the viable cell count procedure. For this, at the pre-established periods a 100  $\mu$ L aliquot of the media was uniformly spread on sterile SSDL agar Petri dishes and incubated at 37°C for 48 h. After the incubation period the count of viable cell was carried out and the results were expressed as Log of Colony Forming Units per mL (Log cfu mL<sup>-1</sup>).

Concentration of ethanol was determined using a gas chromatograph (HP 5890, Hewlett-Packard, Palo Alto, CA) fitted to a flame ionizer detector. A 2  $\mu$ L-portion of the fermentation sample was injected onto a column (30 m; 0.25 mm i.d.; 0.25 lm, J&W Scientific, Folsom, CA). The chromatographic conditions were as follow: sample (without dilution) injection volume 2  $\mu$ L; hydrogen flow rate 5.0 mL min<sup>-1</sup>; temperature program 120°C (isotherm); injector temperature 100°C; detector temperature 120°C. The data were processed using the Millennium Computer Program (Waters Chromatograph Division, Milford, MA, USA). Analyses were performed in triplicate and the results were expressed as average of the parallel assays.

The following parameters were used for assessing the fermentative process: (i) Amount of consumed sugar: [S: -  $(S_f - S_o)$ ], where S: consumed sugar (g glucose L<sup>-1</sup>);  $S_o$ : sugar final concentration (g glucose L<sup>-1</sup>);  $S_o$ : sugar initial concentration (g glucose L<sup>-1</sup>); (ii) Amount of produced ethanol: [ $P: (P_f - P_i)$ ], where P: amount of produced ethanol (g L<sup>-1</sup>);  $P_i$ : final ethanol concentration (g L<sup>-1</sup>);  $P_i$ : initial ethanol concentration (g L<sup>-1</sup>);  $P_i$ : not glucose g ethanol<sup>-1</sup>); (iv) Rate of substrate yield to ethanol (g glucose g ethanol<sup>-1</sup>); (iv) Rate of ethanol production: [PR: P/t], where PR: rate of ethanol production (g L<sup>-1</sup> h<sup>-1</sup>); t: time of fermentation (h); (v) Efficiency of fermentation ( $n_{p(N)}$ ), based on the theoretical yield according to the Gay-Lussac equation (51.1 g ethanol 100 g glucose<sup>-1</sup>): [ $n_{p(N)}$ ;  $Y_{p/s}$ .100/51.1].

The data were analyzed by ANOVA using the Duncan test ( $p \le 0.05$ ) and the software Statistica 6.0.

# Results

During the fermentation the amount of glucose increased up to 24 h (Figure 1). After 30 h of fermentation the glucose concentration dropped sharply and it was absent in the media after 40 h of fermentation. Along the 72 h fermentation period, the pH value of mesquite pods mash did not vary (4.8 - 5.1). The maximum ethanol amount (165 g L<sup>-1</sup>) was achieved at 36 h of fermentation (Figure 1). The highest experimental count of Z. mobilis (close to 6 Log cfu mL<sup>-1</sup>) was also noted at 36 h (Figure 1). Highest rates of substrate yield to ethanol (0.44 g ethanol g glucose<sup>-1</sup>) and efficiency of fermentation (86.81%) were found after 30 h (Table 1). Ethanol productivities were 4.69 g L<sup>-1</sup> h<sup>-1</sup> and 4.56 g L<sup>-1</sup> h<sup>-1</sup> noted after 30 h and 36 h of fermentation, respectively (Figure 1). At 72 h the values found for all assessing parameters dropped sharply. 30 h was found as the shorter interval time to obtain the highest ethanol yield from the fermentation of mesquite pods mash by Z. mobilis under static condition.

#### Discussion

The amount of glucose dispersed in the growth medium was increasing up to 24 h of fermentation, although it was suppressed after 40 h. The increasing glucose amount found in mesquite mash up to 24 h was probably related to a continuous hydrolysis of sucrose dispersed in the medium. Silva et al. (2007) reported that mesquite pods present high availability of sugars (56.5 g

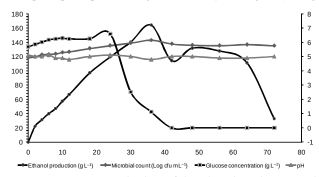


Figure 1 – Experimental values of the ethanol production and microbial count in submerged fermentation of hydrat mesquite pods mash by Z. mobilis UFEPEDA-205 under static condition.

100 g<sup>-1</sup>), particularly sucrose. High availability of sucrose in the growth media increases the ethanol yield by *Z*. *mobilis* (Favela Torres and Barati, 1988). On the other hand, small yields of ethanol are found in media rich of cellulose, inulin or starch since the bacterium is not able to hydrolyze these polymers (Shene and Bravo, 2001).

Z. mobilis presents a prominent capacity of hydrolyzing the sucrose dispersed in the growth media and rapidly metabolizes the resulting glucose as carbon source for ethanol production by the Entner-Doudoroff way (Swings and De Ley, 1977). The availability of glucose, fructose or sucrose in the growth media increases the ethanol yield by Z. mobilis (Favela Torres and Barati, 1988). Extremely high levels of glucose in a basal medium could suppress the growth of Z. mobilis while causing no negative effect toward the ethanol yield (Morais et al., 1993). The non-occurrence of changes at stabilization of pH in the mesquite mash during 72 h of fermentation could indicate the absence of contaminating bacteria in high number during the fermentation (Narenddranath and Power, 2004).

The three times of fermentation (30, 36 and 72 h) evaluated were chosen based on the higher and smaller obtained ethanol yield. The findings suggested 30 h as the shorter interval time to get the highest ethanol yield from the fermentation of mesquite pods mash by Z. *mobilis* under static condition, which could possibly reduce the overall time of the process and decrease the cost for industrialization.

Aerobic cultures of Z. mobilis had higher ethanol yield from glucose with maximum theoretical values of 0.51 g ethanol g glucose<sup>-1</sup> (Prasad et al., 2007). On the other hand, small yields of ethanol from glucose (0.13 to 0.17 g ethanol g glucose<sup>-1</sup>) are found in anaerobic cultures (Viikari, 1998; Shene and Bravo, 2001). Tano and Buzato (2003) reported a small ethanol production (29 g L<sup>-1</sup>) from sugar cane juice by Z. mobilis ATCC 31821 with a small substrate yield into ethanol (0.42 g ethanol g glucose<sup>-1</sup>) after 48 h under stirring (aerobic) cultivation. According to the authors, this small yield was possibly related to the amount of mineral compounds dispersed in the sugar cane juice that could inhibit the fermentation by Z. mobilis. Regarding the results of previous studies the mesquite pods hydrated mash present a higher efficiency for ethanol production in comparison to the classical substrate sugar cane juice.

Table 1 – Assessing parameters of the fermentative process for ethanol production from mesquite pods hydrated mash by *Z. mobilis* UFEPEDA-205 under static conditions for 72 h.

Assessing parameters	Time of fermentation (h <sup>-1</sup> )		
	30	36	72
$Y_{p/s} (g g^{-1})$	0.44 ± 0.02 a	0.35 ± 0.03 b	0.05 ± 0.02 c
PR (g L <sup>-1</sup> h <sup>-1</sup> )	$4.69 \pm 0.02$ a	4.56 ± 0.04 b	0.45 $\pm$ 0.01 c
$N_{p(\%)}$ (%)	86.81 ± 0.02 a	70.37 ± 0.01 b	11.27 ± 0.02 c

Average values followed for different letter at the lines differ (Duncan, p d" 0.05);  $Y_{p/s}$  rate of substrate to ethanol conversion; PR: ethanol productivity;  $N_{p/s}$ : efficiency of fermentation

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Our findings suggest mesquite pods as an emerging substrate for ethanol production using submerged fermentation by Z. *mobilis*. High levels of ethanol were produced from mesquite pods hydrated mash by submerged fermentation using Z. *mobilis* under static condition.

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