Interaction of Lactobacillus vini with the ethanol-producing yeasts Dekkera bruxellensis and Saccharomyces cerevisiae

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Abstract

Lactobacillus vini was recently described as a contaminant in industrial ethanol fermentations and its co-occurrence with Dekkera bruxellensis was noted. We investigated the growth characteristics of L. vini in cocultivation together with either Saccharomyces cerevisiae or D. bruxellensis. Lower cell numbers of both the yeasts and L. vini as well as a decrease in ethanol and lactate formation in mixed batch cultures compared with pure cultures were noted. L. vini formed cell aggregates (flocs) in all cultivation media with different shapes in Man–Rogosa–Sharpe and yeast extract–peptone–dextrose media. Flocs’ size and proportion of cells bound to flocs increased with increasing ethanol concentration. In coculture, formation of lactic acid bacteria–yeast cell aggregates consisting of a bacterial core with an outer layer of yeast cells was observed. L. vini–D. bruxellensis flocs had a bigger surface, due to cells protruding from the pseudomycelium. The involvement of mannose residues in the flocculation between L. vini and yeasts was tested. The presence of mannose induced deflocculation in a concentration-dependent manner. Less mannose was required for the deflocculation of D. bruxellensis as compared with S. cerevisiae. © 2013 The Authors. Biotechnology and Applied Biochemistry published by Wiley Periodicals, Inc. on behalf of the International Union of Biochemistry and Molecular Biology, Inc. Volume 61, Number 1, Pages 40–44, 2014

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1. Introduction

Lactic acid bacteria (LAB) constitute one major group of contaminants in industrial yeast fermentations, and a variety of species have been isolated from those processes [1–3]. Lactobacillus fermentum and Lactobacillus brevis were reported to be the most common contaminants in ethanol production [4]. High growth rates, tolerance to low pH and high temperature, and resistance to high ethanol concentrations enable a variety of LAB to compete directly with yeast for the sugar substrate and may also inhibit the fermentation process by producing organic acids, mainly lactic acid. The species Lactobacillus vini was described a few years ago [5]. The bacterium has been isolated from several ethanol production plants in Sweden and Brazil; therefore, there is a demand to understand the interaction of L. vini with yeasts [2, 6]. The involved ethanol plants run in a continuous mode with cell recirculation, at sugar limitation and ethanol concentrations up to 100 g/L. In the Swedish plant, low pH (3.6) and temperatures of 36 °C have been documented [6]. In most documented cases, L. vini was associated with the presence of Dekkera bruxellensis. D. bruxellensis is a nonconventional yeast that outcompeted traditionally used Saccharomyces cerevisiae in several ethanol production plants running continuous fermentation with cell recirculation [2, 6]. Although D. bruxellensis is usually considered as a contaminant in ethanol production [7, 8], it has been shown to have ethanol yields similar to or even higher than S. cerevisiae as well as low glycerol formation, high biomass yield, a high ethanol tolerance, and a broad range of temperature and pH tolerance [9–11]. On the other hand, it has lower specific ethanol production and growth rates than S. cerevisiae and may therefore not be competitive in batch fermentations.
It is currently unclear why *D. bruxellensis* so frequently co-occurs with *L. vini* in industrial ethanol fermentation [12]. To investigate LAB–yeast interactions, cocultivations under conditions that are conducive to both microorganisms are required. This study aimed to test growth requirements of *L. vini* and to assess its growth dynamics in batch pure cultures and in cocultures with *D. bruxellensis* and *S. cerevisiae*.

2. Materials and Methods

2.1. Strains and culture media

The yeasts strains *D. bruxellensis* CBS 11270 and *S. cerevisiae* J672 [9] and the LAB strain *L. vini* 2FLAB5 isolated from a Swedish ethanol plant [6] were used in this study. *L. vini* was pregrown on Man–Rogosa–Sharpe (MRS) agar (Oxoid, Basingstoke, New Hampshire, England) in an anaerobic environment using a BD BBL™ GasPak™ Plus Anaerobic System (Becton, Dickinson and Company, Franklin Lakes, NJ, USA). Yeast extract–peptone–dextrose medium (YPD) contained 10 g/L yeast extract, 20 g/L bacteriological peptone (both from Oxoid), and 20 g/L d(+)-glucose (VWR International Ltd., Poole, England). Solid media also contained 16 g/L agar. For testing the influence of ethanol on flocculation, 10, 50, or 150 g/L ethanol were added to liquid YPD. Synthetic medium contained KH$_2$PO$_4$ 18.75 g/L, (NH$_4$)$_2$HPO$_4$ 6 g/L, MgSO$_4$ 7H$_2$O 1.13 g/L, yeast nitrogen base (Difco™; Becton Dickinson and Company) 6.5 g/L, and glucose 40 g/L. Several supplements of synthetic medium were tested: 5 g/L yeast extract or a mixture of ergosterol (18 µg/L) and Tween 80 (3 g/L) combined with 20 g/L bacteriological peptone or all 20 proteinogenic amino acids (100 mg/L each). The ergosterol/Tween 80 solution was prepared according to Ref. [13].

2.2. Precultures

Yeast precultures were incubated in 150 mL synthetic medium in 500 mL shake-flasks with shaking (200 rpm) at 30 °C for approximately 72 h for *D. bruxellensis* and for 24 h for *S. cerevisiae*.

Precultures of *L. vini* were grown in 200 mL MRS medium [MRS broth 52 g/L, (Oxoid)] in 250 mL flasks (Schott Duran, Wertheim/Main, Germany) with shaking (200 rpm) at 30 °C for approximately 100 h.

2.3. Growth tests of *L. vini*

*L. vini* was inoculated to an OD$_{600}$ of about 1.0 (determination was not exact because flocs could not completely be resuspended by vigorous shaking) into 20 mL serum vials (PharmaPack Packaging System AB, Södertälje, Sweden) containing 10 mL of corresponding medium. The initial pH was adjusted to 5, using concentrated hydrochloric acid. Experiments were performed in triplicate. The vials were incubated at 30 °C and shaken at a speed of 200 rpm on a rotary table.

2.4. Batch cocultivation

For cocultures, every tested microorganism was inoculated to an OD$_{600}$ of approximately 1 into 20 mL serum vials containing 15 mL of YPD, which were closed with a rubber stopper perforated with a syringe needle. The initial pH was adjusted to 5. Pure cultures inoculated to an OD$_{600}$ of approximately 1 were run as controls. Experiments were performed in triplicate. The vials were incubated at 30 °C and shaken at a speed of 200 rpm on a rotary table.

2.5. Influence of mannose on flocculation

Precultured yeast and bacterial cells were washed with sterile saline (NaCl solution, 9 g/L). For testing the effect of mannose on LAB and LAB–yeast flocculations, every examined microorganism was inoculated to an OD$_{600}$ of approximately 0.2 into 2 mL tubes containing 2 mL of 10$^{-1}$, 10$^{-2}$, 10$^{-3}$, 10$^{-4}$, or 10$^{-5}$ M mannose in a buffer solution (9.35 g/L KH$_2$PO$_4$, 3 g/L (NH$_4$)$_2$PO$_4$, 1 g/L MgSO$_4$ × 7H$_2$O, pH 5). Experiments were performed in triplicate.

2.6. Analytical methods

Ethanol, acetate, and glycerol were measured by high-pressure liquid chromatography as described earlier [14]. Cell viability was determined by viable counts where serial dilutions were plated on YPD-agar plates. The plates were incubated at 30 °C for 24 h (*S. cerevisiae*) or 72 h (*D. bruxellensis*). *L. vini* plates were kept in anaerobic jars. Differentiations of the yeasts and bacteria species were based on the differences in colony morphology. *S. cerevisiae* colonies had a yellowish color and dull surface with deepening in the center. *D. bruxellensis* had smaller colonies of a whitish color with a glittering and protuberant surface. *L. vini* colonies were the smallest size with a swelling center and had a whitish translucent color.

Flocculation was studied at 400-fold magnification using an Olympus BH2 microscope. The influence of mannose on flocculation was also monitored by optical density measurements immediately after suspending cells in the buffer and 7 Min after at a wavelength of 600 nm (OD$_{600}$) by an Ultrospec 1100 pro spectrophotometer.

2.7. Statistical analysis

To test whether there were significant differences between the treatments, Student’s *t*-test was performed with a significance level of 5% using the software MS Excel (Version 2007).

3. Results

3.1. Growth characteristics of *L. vini* in batch culture

We initially determined the growth of *L. vini* in different media that could potentially be used for cocultivation of LAB and yeasts. *L. vini* was able to grow in YPD medium, reached approximately 1.0 × 10$^8$ cells/mL (Fig. 1), and did not increase further even though there was still a significant amount of glucose present in the medium (8.4 ± 1.1 g/L).

We tested whether it was possible to replace components in the rich medium with defined supplements. Yeast extract could be replaced by a mixture of ergosterol and Tween 80, which is also essential for anaerobic cultivation of yeasts [15].
However, peptone could not be replaced by a mixture of all 20 amino acids. It is possible that \textit{L. vini} requires oligopeptides for growth, which has been observed in other LAB species \cite{16}. Substantial flocculation was observed in all media tested in this study. Flocs in YPD medium were spherical, whereas flocs in MRS medium tended to be irregular clumps. When ethanol was added to the medium at concentrations of 10, 20, and 100 g/L, the degree of \textit{L. vini} flocculation increased in a concentration-dependent manner. We observed both cell aggregates of different sizes as well as single cells in the absence of ethanol. The amount of single cells decreased when 10 g/L ethanol was added to a growing culture, and no individual cells could be detected at 100 g/L ethanol.

3.2. Batch cocultivation of \textit{L. vini} with yeast

To investigate the impact of \textit{L. vini} on yeast growth, it was cocultivated with \textit{D. bruxellensis} and \textit{S. cerevisiae}. In both cocultures with \textit{S. cerevisiae} and \textit{D. bruxellensis}, the cfu number of \textit{L. vini} reached the same level of $7.1 \pm 2.0 \times 10^6$ cfu/mL, which was only 70\% of the corresponding cell number in pure culture (Figs. 2A and 2B). Lactate production was significantly reduced by 10\%–20\% in both cocultures. In coculture, \textit{D. bruxellensis} viable counts were higher than those of \textit{S. cerevisiae} ($5.0 \times 10^7 \pm 1.5 \times 10^6$ cfu/mL compared with $4.5 \times 10^7 \pm 1.5 \times 10^6$ cfu/mL), respectively. However, the final ethanol concentration in \textit{L. vini–D. bruxellensis} coculture was significantly ($P = 0.016$) lower (5.3 \pm 0.2 g/L) than in the \textit{L. vini–S. cerevisiae} coculture (6.0 \pm 0.2 g/L). Final ethanol concentrations in the corresponding \textit{D. bruxellensis} and \textit{S. cerevisiae} pure cultures were 7.2 \pm 0.1 and 7.3 \pm 0.4 g/L, respectively (Fig. 2). Low amounts of glycerol and acetate were also formed in both cocultures. The maximum concentrations after glucose was finished for glycerol and acetate, respectively, were $0.04 \pm 0.03$ and $0.19 \pm 0.08$ g/L in the \textit{D. bruxellensis–L. vini} coculture and $0.52 \pm 0.02$ and $0.19 \pm 0.05$ g/L in the \textit{S. cerevisiae–L. vini} coculture.

3.3. Coflocculation of \textit{L. vini} and yeast

Microscopic investigation of \textit{L. vini–yeast} cocultures revealed that \textit{L. vini} formed aggregates together with either yeast (Figs. 3A and 3B). Cell aggregation occurred immediately after addition of the \textit{L. vini} cells to the yeast cultures. The flocs consisted of a bacterial core surrounded by yeast cells. Several aggregates were completely covered with yeast cells. Moreover, cells of \textit{D. bruxellensis} formed pseudohyphae that protruded from the flocs of budding cells of \textit{S. cerevisiae} mainly formed a monolayer around the bacterial core (Figs. 3A and 3B).

3.4. Influence of mannose on flocculation

The degree of \textit{L. vini, L. vini–D. bruxellensis}, and \textit{L. vini–S. cerevisiae} flocculation decreased when mannose was present in the suspension in a concentration-dependent manner. This effect was more pronounced 7 Min after suspending the cells in the buffer as compared with the time zero (Fig. 4). This was due to sedimentation of the flocs in samples with low mannose content. Microscopic investigation revealed a difference in the mannose effect on the \textit{L. vini–D. bruxellensis} and \textit{L. vini–S. cerevisiae} flocs. The concentration efficient for complete deflocculation of \textit{L. vini–D. bruxellensis} was $10^{-2}$ M mannose, whereas it was $10^{-1}$ M for \textit{L. vini–S. cerevisiae}. Mannose addition also affected floc formation by bacterial cells both in pure and in mixed culture. However, LAB flocs were never completely resolved within the tested mannose concentrations.
4. Discussion

This study investigated the growth of the ethanol-tolerant LAB *L. vini* in liquid medium that can be used for cocultivation with yeasts. A high degree of flocculation was observed for this bacterium in all tested media. A similar phenomenon was shown for other LAB species [17, 18]. Possibly, the formation of flocs can reduce the surface exposed to ethanol and thus help the bacteria to survive the stressful conditions of the ethanol process. This conclusion is also supported by the observed higher flocculation activity of *L. vini* in the presence of ethanol.

On the other hand, a smaller surface area also decreases sugar and nutrient uptake by the cells.

Interestingly, *L. vini* formed coaggregates with the yeast cells when cultivated in mixed cultures. Coflocculation between yeasts and LAB has rarely been investigated, but it has been shown that *L. fermentum*, which is also present in industrial ethanol processes, can also form aggregates with *S. cerevisiae* [19]. The tendency of *L. vini* to form aggregates with the yeast makes it difficult to detect contamination by microscopic investigations when the bacteria are hidden in the flocs.

A growth cessation of *L. vini* in pure culture was observed while glucose was still present in the medium. The likely reason for this should be the relatively high concentration of lactate (10–12 g/L). The LAB *L. fermentum* and *L. brevis* also stopped growing in the presence of 10 g/L of lactate [4].

In industrial ethanol processes with cell recirculation, *L. vini* was at several occasions demonstrated to be associated with *D. bruxellensis* [2, 6]. The cocultivation experiments described in this study are a first attempt to understand the specific interaction between these two species; however, the results of our experiments did not provide an obvious explanation for this frequent coexistence. *D. bruxellensis* seemed to be more affected by *L. vini* than *S. cerevisiae*, because ethanol concentration decreased on the expanse of formed lactic acid in the corresponding coculture. However, this may apply only for a batch culture, where the conditions are not favorable for
due to its lower specific glucose uptake and growth rates [9]. The behavior during the stationary phase might closely resemble the conditions in the continuous ethanol process, but no impact on survival during this phase was seen in any of yeasts.

In the batch culture, L. vini apparently acts as a contaminant, as it channels sugar to lactate and bacterial biomass, at the expense of ethanol formation and yeast cell numbers. There is no proof yet about whether and how L. vini influences the competition between D. bruxellensis and S. cerevisiae. Attachment of LAB to yeast cells is attributed to mannose binding proteins of the bacteria [20]. Observed differences in the mannose effect on the L. vini–D. bruxellensis and L. vini–S. cerevisiae flocs may be due to the higher mannose content of the cell wall of S. cerevisiae, compared with D. bruxellensis [21], which can lead to higher capacity for L. vini binding to S. cerevisiae. Cells incorporated within the flocs tend to sediment while single cells stay in suspension. Thus, frequent cooccurrence of D. bruxellensis and L. vini may be due to the fact that D. bruxellensis is less involved in flocculation and thus less impacted by L. vini.

It has been found that D. bruxellensis can outcompete S. cerevisiae in glucose limited cocultivations without the involvement of any bacteria [22]. However, it is possible that the smaller flocs with the protruding cells formed by D. bruxellensis–L. vini represent an advantage for the cells because the surface is enlarged, enabling more efficient uptake of the limited nutrients while at the same time protecting the cells in the inner part of the aggregate from stressful conditions [23, 24]. Thus, our results indicate that LAB can affect ethanol production not only by competing for substrate or lactic acid formation but also by influencing spatial structure of the population in the fermentor.

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