Carrier-free, continuous primary beer fermentation

Eduardo J. Pires,1* José A. Teixeira,1 Tomás Brányik2 and António A. Vicente1

Developing a sustainable continuous fermentation reactor is one of the most ambitious tasks in brewing science, but it could bring great benefits regarding volumetric productivity to modern breweries. Immobilized cell technology is often applied to reach the large densities of yeast needed in a continuous fermentation process. However, the financial cost associated with the use of carriers for yeast immobilization is one of the major drawbacks in the technology. This work suggests that yeast flocculation could address biomass immobilization in a gas-lift reactor for the continuous primary fermentation of beer. Nearly 25 g dry wt L−1 of yeast was flocculated in the reactor before interruption of the fermentation. Stable sugar consumption and ethanol production (4.5% alcohol by volume) from an 11°P wort was evidenced. The key esters and higher alcohols measured in the young beer met the standards of a finished primary beer fermentation. Copyright © 2014 The Institute of Brewing & Distilling

Keywords: continuous beer fermentation; yeast flocculation; gas-lift reactor

Introduction

Continuous beer fermentation is far from being a recent science, with patent registrations dating back the nineteenth century. Indeed, the first continuous bioreactors were designed to work with freely suspended/flocculated yeast. However, many factors hindered the evolution of ‘non-immobilized’ continuous systems in the late 1970s, such as: inflexibility in production rate and beer type; lack of control over wort attenuation; excess ancillary equipment needed; difficulty in maintaining production scale hygiene; yeast mutations; and wild yeast contamination (1). The idea of using solid carriers came later to overcome the pitfalls presented by ‘non-immobilized’ reactors, mainly (but not limited to) how to increase the productivity rate beyond the washout rate of yeast. Immobilized cell technology (ICT) uses solid carriers for cell adsorption and biofilm formation (2–9) and forced physical entrapment of yeast cells to solid matrices (10–13) as methods to increase cell density inside the reactor. The carrier cost is a key component for the financial viability of ICT (1,14) and the composition of the solid matrix may also interfere in the final beer quality and flavour profile (15,16). Additionally, the relatively short lifespan of a single yeast cell (17) results in an accumulation of dead biomass in the biocatalyst, thus demanding constant replacement (18).

Continuous beer fermentation depends on a high density of yeast cells immobilized inside a bioreactor. The considerable amount of yeast allows a short residence time for the wort in the bioreactor. The wort is continuously supplied for beer production. This feature results in high productivity and reduced space and time being needed to reach the final product (15,19,20). Despite being very attractive, the continuous fermentation of beer is still marginalized to laboratory benches, pilot plants and a few courageous companies such as Dominion Breweries of New Zealand, which have been using continuous brewing since 1959 with a setup based on flocculated biomass (15).

Some brewing yeast strains are remarkably flocculent and this feature is widely used to harvest yeast at the end of primary fermentation (21,22). Flocculation is a multifactorial inheritance triggered by both genetic and environmental factors (22). The most accepted mechanism supporting yeast flocculation is through the expression of FLO genes encoding specialized cell wall proteins (flocculins), which are able to bind to sugar residues in the cell walls of adjacent cells (22,23). Hydrodynamic conditions in the reactor have a direct impact on flocculation and floc size, as the liquid agitation increases the chances of cell collisions, but strong movement breaks up cell clusters (24). Moreover, the higher the concentration of yeast cells that are in suspension, the greater the number of collisions, and consequently the faster the formation of flocs (25). Additionally, factors that increase cell-surface hydrophobicity and/or that decrease the repulsive negative electrostatic charges on the cell wall cause stronger flocculation, as they increase the probability of cell-to-cell contact (26). This self-aggregation characteristic is a free-of-charge way of immobilization and seems to have been poorly exploited in recent ICT setups.

Gas-lift bioreactors are remarkably efficient in regard to mass transfer for either liquid–liquid or solid–liquid phases triggered by rapid mixing, low shear stress, simple design and low energy consumption (27,28). The low shear environment provides excellent conditions for ICT implementation (29) and has previously been adopted for continuous beer fermentation (3,13). This work examines the feasibility of self-aggregation as a single biomass immobilization method for young beer production in a gas-lift reactor.

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Material and methods

Brewing yeast

The flocculent Saccharomyces pastorianus strain no. 96 from the Culture Collection of Brewing Yeast (Research Institute of Brewing and Malting plc, Prague, Czech Republic) was used. It was inoculated onto a complex media of the following composition (g L⁻¹): glucose, 30; KH₂PO₄, 5; MgSO₄·7H₂O, 0.4; (NH₄)₂SO₄, 2; yeast extract, 2; agar, 20. The medium was prepared in Petri dishes for the isolation of yeast colonies. A colony was then inoculated in 400 mL of 5°Plato wort and incubated at 20°C, 120 rpm, for 48 h before use as an inoculum for the gas-lift reactor.

Beer wort

Pale wort concentrate was acquired from the Research Institute of Brewing and Malting (Prague). It was diluted to the final desired concentration – 5° and 11°P. The final wort was filled into 20 L polyethylene carboys (Nalgene, USA) and autoclaved for 3.5 h.

Continuous fermentation

The experiment was carried out using a Perspex gas-lift reactor with 4 L of total work volume. Gas flow was kept constant at 0.5 L min⁻¹ by the GFC17 mass controller (Aalborg, USA). The temperature was held at 15°C using a Julabo F32 refrigeration/heating circulator (Julabo, Germany). The dilution rate was kept at 0.043 h⁻¹ using a peristaltic pump PDC 83 (Kou, Czech Republic). The reactor was sterilized by bleaching, using a 3% (v/v) solution of commercial sodium hypochlorite with 1.5% of active chlorine, 48 h prior to use. After this time, the solution was discarded and 50 L of sterile water was used to wash the reactor. It was then filled with 5°P wort and inoculated.

Batch growth was performed in the first 48 h, using pressurized air as the gas supply at 500 mL min⁻¹. After that, the gas was changed to CO₂ at the same flow rate and the continuous phase was started with a 5°P wort supply at a 0.043 h⁻¹ dilution rate. Four days later, the wort supply was changed to 11°P wort keeping the dilution rate unchanged.

Biomass measurements

The flocculated biomass was evaluated daily. Three 15 mL plastic falcon tubes were dried at 105°C for 12 h and weighed. Then 10 mL of cell suspension from the reactor was added to each one of them and centrifuged at 4000 g for 5 min. The liquid phase was discarded and the tubes were dried (105°C) for 24 h prior to weighing. Control blank experiments were carried out using the inlet wort to correct for the presence of turb-like components that could interfere with weighing.

Sugars and ethanol measurements

A daily sample was taken from the reactor’s outflow for green beer analysis. Sugars and ethanol were evaluated using high-performance liquid chromatography (HPLC) with an Agilent 1100 series equipped with Agilent G 1362A RID detector (Agilent, USA). The column used was the Rezex™ RS-O-Oligosaccharide 200 × 10 mm (Phenomenex, USA) and the eluent was deionized degassed water pumped at a flow of 0.4 mL/min. Sugar and ethanol standards were calibrated previous to wort and young beer measurements using the following reagents: d-fructose (Chemapol, Praha, Czech Republic), d-maltose monohydrate (Fluka, Japan), d-glucose (Fluka, Japan), maltotriose (Sigma, USA) and ethanol (Sigma, USA).

Measurement of flavour-active compounds

Higher alcohols and esters were analysed by gas chromatography using an Agilent HP-6890N gas chromatography–mass spectrometry system (Agilent Technologies, USA) coupled to a mass detector Agilent 5975B Inert MSD (Agilent Technologies, USA). Compounds were separated on an Innowax (30 m × 0.25 mm × 0.25 μm) column (Agilent Technologies, USA). Helium was used as the carrier gas at a flow rate of 6 mL/min. The oven temperature was programmed to a start temperature of 30°C for 10 min, then it was raised at 2°C min⁻¹ to 52°C (2 min), plus 2°C min⁻¹ up to 65°C and finally up to 250°C at 5°C min⁻¹ (3 min). Samples were injected at 260°C. Standards were analysed previous to samples using 2-methyl-1-butanol (>98%), 3-methyl-1-butanol (>98.5%), isobutanol (>99%), isomyl acetate (<99%), ethyl acetate (99.7%), ethyl butyrate (>98%), ethyl hexanoate (>99%), ethyl octanoate (>98%), ethyl decanoate (>99%) and 2-phenyl ethyl acetate (>99%) (Fluka, Germany). Internal standards used were 3-octanol (99%; Aldrich, USA) and ethyl heptanoate (99%; Aldrich, Germany).

Results and discussion

Biomass build-up

One of the main conditions in the environment that triggers yeast flocculation is the reduced amount of sugars present at the end of primary fermentation (22,30,31). For this reason diluted (5°P) wort was used in the start-up of the gas-lift reactor. Unexpectedly, this lower input of sugars did not favour biomass
was virtually impossible owing to the small size of the flocks and the dynamics of cell aggregation (38,39). Hence, the biomass expressed in Fig. 2 denotes the total biomass ($X_{\text{tot}} = \text{flocs} + \text{free cells}$) suspended in the system.

Comparing $X_{\text{tot}}$ in different setups used by other authors is a difficult task. Whereas cell adsorption is very dynamic, entrapment-based studies often do not discuss biomass growth on biocatalysts. For example, Tata and co-workers (8) compared a number of two-stage systems for continuous beer production. Each of these systems was composed of two reactors connected in series: two fluidized bed reactors with porous glass beads for cell immobilization and two loop reactors with a silicon carbide cartridge for yeast load. The maximum values of $X_{\text{tot}}$ reported by the authors for each of these systems were 29.7 and 18.2 g cell dry wt L$^{-1}$, respectively. In further work involving continuous primary beer fermentation carried out by Brányik et al. (40), the $X_{\text{tot}}$ reported using a gas-lift reactor with a lignocellulosic carrier obtained from brewer’s spent grains varied from 9.3 to 10.5 g cell dry wt L$^{-1}$. However, the yeast load of a specific system does not necessarily reflect good results. Therefore, the performance of different ICT setups must be compared in terms of specific saccharide consumption ($r_s$) and volumetric ethanol productivity ($r_p$) (8,41) as further discussed.

** Sugars consumption and ethanol yield **

The efficacy of the gas-lift reactor for primary beer fermentation has long been studied, using several types of yeast immobilization method (13,40,42). The present work focused on self-aggregation to attain the biomass needed for the continuous primary fermentation of the wort provided. As temperature and dilution rate were kept constant during the entire experiment, sugar consumption and ethanol production were directly dependent on the $X_{\text{tot}}$ and wort gravity. Table 1 shows in detail the amount of each fermentable sugar present at the inlet wort for both the 5 and the 11°P stages. It also shows the residual content of these sugars and the ethanol present in the green beer through the course of the continuous fermentation. Changing the composition of wort during the continuous phase caused an adaption phase between the 144 h (start of inlet supply of 11°P barrel) and 388 h (beginning of steady phase) period of the continuous fermentation (Fig. 2). Environmental changes are known to alter yeast metabolism,

![Figure 2](image-url)

**Figure 2.** Specific saccharide load and consumption ($r_s$), ethanol productivity ($r_p$) and total immobilized biomass ($X_{\text{tot}}$) inside the reactor through the primary continuous fermentation. This figure is available in colour online at wileyonlinelibrary.com/journal/brewing.
which can then take some time to adapt to the new conditions \((43)\). Thus, instead of changing the wort gravity, the dilution rate should have been raised gradually, using a steady wort gravity, to avoid delays. After the adaption stage, the consumption of sugars was satisfactory and stable until the end of the fermentation trial. At this time, from the original \(84.25 \text{ g L}^{-1}\) of fermentable sugars present in the inlet \((11^\circ P)\) wort, only about \(5 \text{ g L}^{-1}\) was present in the outflow. These residual sugars corresponded to maltose and maltotriose (on average 2 and 3 g L\(^{-1}\), respectively). Not surprisingly, glucose and fructose were not present in the young beer at this phase \((43)\) owing to preferential sugar consumption by yeast. The threshold for these alcohols when considered together is \(50 \text{ mg L}^{-1}\) and this value increased during the steady phase of \(11^\circ P\) wort supply, the total retention time measured during this period \((44)\). As the total retention time \((R_{\text{tot}})\) was 0.66 g L\(^{-1}\) of amyl alcohol content. Higher alcohols not only contribute directly to the final beer aroma, but can also be used as precursors in the synthesis of esters. Data involving the production of higher alcohols using ICT are rather inconsistent as some works have reported a low production of higher alcohols using ICT \((51–53)\), while others have observed even higher values of fusel alcohols in continuous mode when compared with batch fermentation \((54)\). In the current work, isoamyl alcohol \((3\text{-methyl-1-butanol})\) and the active amyl alcohol \((2\text{-methyl-1-butanol})\) were measured together and are henceforth reported as amyl alcohols. The threshold for these alcohols when considered together is \(50 \text{ mg L}^{-1}\) \((55)\). Higher alcohols measured in the green beer produced in the current work are presented in Table 2. When fermentation reached a steady stage upon \(11^\circ P\) wort supply, the threshold of amyl alcohols was achieved. At this time, the \(X_{\text{tot}}\) was nearly 14 g dry wt L\(^{-1}\) and the amyl alcohol concentration was 52 mg L\(^{-1}\). Levels of these alcohols changed slightly \((55 \text{ mg L}^{-1})\) up until the interruption of fermentation when \(X_{\text{tot}}\) was close to 25 g dry wt L\(^{-1}\). Using polyvinyl alcohol for yeast entrapment, Smogrovicová and co-workers \((56)\) observed 57.8 mg L\(^{-1}\) of amyl alcohol in their young beer. The amyl alcohol yield measured in the Brányik et al. \((47)\) setup using brewer’s spent grains as yeast carriers was somewhat higher and 60.1 mg L\(^{-1}\) was measured. In an early study, Smogrovicová and Dömény used calcium-pectate beads for yeast immobilization and observed a similar amyl alcohol content of 56.8 mg L\(^{-1}\) \((52)\). In this particular last mentioned work, the authors used the same yeast strain and temperature \((15^\circ C)\) and a similar wort gravity \((11.5^\circ P)\) as used in the current work, which may also explain the similarity in the amyl alcohol content. Higher

### Table 1. Wort and green beer composition measured in the course of both \(5^\circ P\) and \(11^\circ P\) wort supply through the continuous primary fermentation

<table>
<thead>
<tr>
<th>Sample type</th>
<th>Time (h)</th>
<th>Maltotriose ((\text{g L}^{-1}))</th>
<th>Maltose ((\text{g L}^{-1}))</th>
<th>Glucose ((\text{g L}^{-1}))</th>
<th>Fructose ((\text{g L}^{-1}))</th>
<th>ABV ((%))</th>
<th>Ethanol ((\text{g L}^{-1}))</th>
<th>(X_{\text{tot}}) ((\text{g dry wt L}^{-1}))</th>
</tr>
</thead>
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<tr>
<td>(5^\circ P) wort</td>
<td>—</td>
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<td>23.49</td>
<td>5.71</td>
<td>2.46</td>
<td>—</td>
<td>—</td>
<td>—</td>
</tr>
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<td>2.34</td>
<td>1.76</td>
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<td>0.32</td>
<td>1.96</td>
<td>15.50</td>
<td>3.9</td>
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<td>13.69</td>
<td>50.96</td>
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<td>6.69</td>
<td>—</td>
<td>—</td>
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<td>20.18</td>
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<td>20.81</td>
<td>5.8</td>
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<tr>
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<td>0</td>
<td>0</td>
<td>4.62</td>
<td>36.54</td>
<td>24.8</td>
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<tr>
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<td>15.57</td>
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</table>

ABV, Alcohol By Volume; GB5, Green Beer from \(5^\circ P\) Wort; GB11, Green Beer from \(11^\circ P\) Wort; \(X_{\text{tot}}\), total biomass.
alcohols production is totally dependent on the fermentation conditions adopted and is usually enhanced by conditions that favour yeast growth (43,57). The isobutanol levels measured in the current work are also in agreement with published data for green beer produced by ICT (47,52). Surprisingly, the level of 2-phenylethanol was high in the green beer produced from 11̊P wort (Table 2), which could ultimately contribute to a pleasant floral aroma in the finished beer.

Esters are pleasant aroma compounds present in beer and contributing positive notes such as flowers, honey and fruity. They are mainly produced during the initial phase of primary fermentation, by the action of yeast acyltransferase activities catalysing the condensation reactions between either acetylacyl-CoA and higher alcohols or ethanol. Accordingly, successful primary beer fermentation must produce enough esters that will be present in the final product. The ester most present in beer is the fruit/solvent-like ethyl acetate, which has a flavour threshold around 25–30 mg L⁻¹ (55,58). For the current work, this threshold was reached at the beginning of the steady phase of the continuous experiment (11̊P). From this time onwards, the ethyl acetate values changed only slightly up to 36.7 mg L⁻¹ at the end of the continuous fermentation. This number is higher than any other data found in the literature and is also in accordance with past works regarding the overproduction of ethyl acetate by ICT systems (3,52,56,59–61). Ester production is greatly influenced by yeast strain, pitching rate, temperature, top pressure, aeration and agitation (43). An increased production of acetate esters by immobilized yeast was evidenced during beer fermentation owing to higher ATF1 gene expression in immobilized cell population when compared with free cells (62). Alcohol acetyltransferase (ATF1 and ATF2) gene expressions are the most important aspects determining acetate ester levels during fermentation (63). Isoamyl acetate is also an important constituent of the final beer with threshold values around 1.2 mg L⁻¹ (55). The performance of the current setup regarding the production of isoamyl acetate was also superior to other data involving primary beer fermentation (3,51,59). However, very high isoamyl acetate levels are not desired in lager beers.

Medium-chain fatty acid (MCFA) ethyl esters such as ethyl hexanoate (caproate), ethyl octanoate (caprylate) and ethyl decanoate (caprate) are also present in beer (Table 2). Among these esters, ethyl hexanoate (caproate) is the most abundant, and its production can be enhanced through the use of immobilized yeast. This improvement was evidenced by higher ATF1 gene expression in immobilized cell population when compared with free cells (62). Alcohol acetyltransferase (ATF1 and ATF2) gene expressions are the most important aspects determining acetate ester levels during fermentation (63).

**Table 2.** Flavour-active compounds present in the outflow (young beer)

<table>
<thead>
<tr>
<th>Compound</th>
<th>Concentration (mg L⁻¹) at:</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>120 h</td>
</tr>
<tr>
<td>Ethyl acetate</td>
<td>3.33</td>
</tr>
<tr>
<td>Isoamyl acetate</td>
<td>0.09</td>
</tr>
<tr>
<td>Ethyl butyrate</td>
<td>–</td>
</tr>
<tr>
<td>Phenyl ethyl acetate</td>
<td>0.30</td>
</tr>
<tr>
<td>Ethyl hexanoate (caproate)</td>
<td>0.008</td>
</tr>
<tr>
<td>Ethyl octanoate (caprylate)</td>
<td>0.005</td>
</tr>
<tr>
<td>Ethyl decanoate (caprate)</td>
<td>0.001</td>
</tr>
<tr>
<td>Total esters</td>
<td>3.73</td>
</tr>
<tr>
<td>Amyl alcohols a</td>
<td>24.55</td>
</tr>
<tr>
<td>Isobutanol</td>
<td>–</td>
</tr>
<tr>
<td>2-Phenylethanol</td>
<td>40.28</td>
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<tr>
<td>Total higher alcohols</td>
<td>64.83</td>
</tr>
<tr>
<td>A/E b</td>
<td>17.38</td>
</tr>
</tbody>
</table>

*a*3-Methyl-1-butanol and 2-methyl-1-butanol; 
*b*higher alcohol to ester ratio.

**Figure 3.** Correlations between the total biomass (Xₜot) present in the gas-lift reactor and the outlet content of total esters and higher alcohols. This figure is available in colour online at wileyonlinelibrary.com/journal/brewing.
decane (caprate) are produced at much lower levels than acetate esters in beer (58). Thus, ICT studies often focus more attention on the acetate family rather than the ethyl ester group. The MCFA ethyl esters measured in this study were consistent with past records involving immobilized yeast during primary beer fermentation (52,56,59–61). Table 2 shows the ester profile of the green beer at different fermentation times for the present work.

Both higher alcohols and esters present in the green beer from the outflow of the gas-lift reactor could be correlated with X_out (Fig. 3) for a specific sampling time. This data can give crucial insights for planning new continuous fermentations. The use of higher pitching yeast rates on the start-up of the reactor may also accelerate the expected results.

Conclusions

The current work has shown that it is feasible to use flocculation as a single method of yeast immobilization in a gas-lift bioreactor for continuous primary beer fermentation. This fact is supported not only by the successful tendency for biomass accumulation relying only on self-aggregation of yeast cells, but also by the good performance on specific saccharide consumption and ethanol volumetric productivity demonstrated in this work. The feasibility of biomass accumulation through flocculation is supported by hydrodynamic (favouring cell–cell collisions) and environmental conditions (low sugar and high ethanol) in the gas-lift reactor. The data obtained was comparable to other ICT systems using several types of yeast carriers in gas-lift reactors. It is important to bear in mind that, although these results are very promising, the experiments were carried out at the laboratory scale and more experiments should be carried out if this setup is to be scaled up. Keeping the composition of the wort as constant as possible is highly recommended as otherwise it could interfere with biomass build-up during the start-up of the reactor. A carrier-free setup would provide advantages concerning the financial cost and management of the reactor when scaled up.

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