Co-fermentation of non-\textit{Saccharomyces} yeasts with \textit{Lactiplantibacillus plantarum} FST 1.7 for the production of non-alcoholic beer

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Abstract
The non-alcoholic beer (NAB) sector has experienced steady growth in recent years, with breweries continuously seeking new ways to fulfil consumer demands. NAB can be produced by limited fermentation of non-\textit{Saccharomyces} yeasts; however, beer produced in this manner is often critiqued for its sweet taste and wort-like off-flavours due to high levels of residual sugars and lack of flavour metabolites. The use of \textit{Lactobacillus} in limited co-fermentation with non-\textit{Saccharomyces} yeasts is a novel approach to produce NABs with varying flavour and aroma characteristics. In this study, lab-scale fermentations of \textit{Lachancea fermentati} KBI 12.1 and \textit{Cyberlindnera subsufficiens} C6.1 with \textit{Lactiplantibacillus plantarum} FST 1.7 were performed and compared to a brewer’s yeast, \textit{Saccharomyces cerevisiae} WLP001. Fermentations were monitored for pH, TTA, extract reduction, alcohol production, and microbial cell count. The final beers were analysed for sugar and organic acid concentration, free amino nitrogen content (FAN), glycerol, and levels of volatile metabolites. The inability of the non-\textit{Saccharomyces} yeasts to utilise maltotriose as an energy source resulted in extended fermentation times compared to \textit{S. cerevisiae} WLP001. Co-fermentation of yeasts with lactic acid bacteria (LAB) resulted in a decreased pH, higher TTA and increased levels of lactic acid in the final beers. The overall acceptability of the NABs produced by co-fermentation was higher than or similar to that of the beers fermented with the yeasts alone, indicating that LAB fermentation did not negatively impact the sensory attributes of the beer. \textit{C. subsufficiens} C6.1 and \textit{L. plantarum} FST 1.7 NAB was characterised as fruity tasting with the significantly higher ester concentrations masking the wort-like flavours resulting from limited fermentation. NAB produced with \textit{L. fermentati} KB112.1 and \textit{L. plantarum} FST1.7 had decreased levels of the undesirable volatile compound diacetyl and was described as ‘fruity’ and ‘acidic’, with the increased sourness masking the sweet, wort-like characteristics of the NAB. Moreover, this NAB was ranked as the most highly acceptable in the sensory evaluation. In conclusion, the limited co-fermentation of non-\textit{Saccharomyces} yeasts with LAB is a promising strategy for the production of NAB.

Keywords Brewing · Non-alcoholic beer · Non-\textit{Saccharomyces} yeasts · Lactic acid bacteria

Introduction

Beer is one of the most widely consumed alcoholic beverages globally; however, despite this, the brewing industry is facing significant challenges. The sector has been severely impacted as a result of the COVID-19 pandemic, with the total volume of beer sold in Europe decreasing by 9% in 2020 [1]. Even prior to this, the beer market had been experiencing a slowdown in overall market growth due to customer demographics, strict legislation, and emerging lifestyle trends [2]. Consumers are becoming increasingly aware of the risks associated with moderate alcohol consumption, such as digestive disorders, cardiovascular disease, cancers, and liver disease [3]; thus, the beer industry is continuously seeking ways to address such consumer demands [2]. Not only does non-alcoholic beer (NAB) allow the consumer to enjoy a beer without the negative effects associated with alcohol consumption, their decreased tax burden and lower production costs offer significant economic opportunities to breweries [4]. In recent years, the global non-alcoholic beer
sector has gained popularity in an ever-increasingly health-conscious population and is forecast to grow at a CAGR of 8.5% by 2026 [5]. From 2013 to 2019, the sold production volume of NAB in the EU increased from 0.59 to 1.38 billion litres, with an estimated value of €1.28 billion. In 2019, over 80% of the volume of NAB produced within the EU could be attributed to five countries; Germany was the top producer accounting for 30.5% of the sold production volume, followed by Spain (16.8%), The Netherlands (14.4%), Poland (12%), and Czechia (7.1%) [6].

NAB may be produced using physical or biological processes. Physical processes involve the removal of ethanol from the final beer through the use of membrane-based or thermal technologies, often resulting in the undesired separation or evaporation of flavour or aroma compounds [7]. Biological processes for NAB production include alteration of the mashing process, cold contact fermentation, limited fermentation, or the use of alternative brewing yeasts [8]. Limited fermentation is the most widely used biological process; however, beer produced in this manner is often criticised for its sweet taste due to residual sugars, aldehyde-associated wort-like off-flavours, and compromised flavour profile due to the limited production of esters [9]. Reviewed by Michel et al. [10] and Bellut et al. [2], another commonly used approach is the use of non-Saccharomyces yeasts which are unable to metabolise the sugars in wort, resulting in decreased ethanol production. Moreover, the application of non-Saccharomyces yeasts which produce high levels of flavour compounds is a promising strategy for the enhancement and optimisation of the flavour profile of NAB produced by limited fermentation. The use of Saccharomyces ludwigii as an alternative brewing yeast has been extensively studied [4, 11–13], while other research has investigated the application of non-conventional yeasts from the genera Candida, Cyberlindnera, Lachancea, Pichia, Hanseniaspora, Torulaspora, Brettanomyces, Schizosaccharomyces, and Zygosaccharomyces, to name but a few [10, 12, 14–18].

Non-alcoholic beers produced by limited fermentation or non-Saccharomyces yeasts often lack the desired pH drop which can lead to microbial spoilage. Traditionally, lactic acid bacteria (LAB) were regarded as beer spoilage microorganisms; however, the presence of LAB in beer fermentation results in the production of organic acids and a subsequent decrease in pH, while also improving the flavour complexity of the beverage through the production of organolectic metabolites, such as alcohols, esters, ketones, aldehydes, and phenolic compounds [19]. Wort acidification by LAB can be performed prior to (pre-fermentation) or in parallel (co-fermentation) with yeast fermentation. LAB pre-fermentation can be carried out in the mashing kettle (mash souring), the brewing kettle (kettle souring) or in the preboiling wort (pre boil wort souring) [19, 20]. Although pre-fermentation eliminates the risk of LAB inhibition by hops or ethanol, the flavour complexity associated with beers produced from mixed fermentations may be lost after boiling [19]. Co-fermentation of LAB and yeast circumvent such an issue, while also shortening beer production times [21]. LAB species which have been investigated for their wort acidification capabilities include Pediococcus acidilactici, Lactiplantibacillus plantarum, Lactobacillus paracasei, Lactobacillus amylovorus, Lactobacillus reuteri, Lactobacillus brevis, Lactobacillus buchneri, and Weisella cibaria [19–25].

The production of NAB by limited co-fermentation of non-Saccharomyces yeasts with LAB has not yet been explored. In this study, we investigated the co-fermentation of Lachancea fermentati KBI 12.1 and Cyberlindnera subsufficiens C6.1 yeasts with L. plantarum FST 1.7 to produce NAB. The commercially available brewer’s yeast Saccharomyces cerevisiae WLP001 was also included as a control. The fermentation kinetics were monitored at time intervals and the composition and metabolite profiles of the final NABs were analysed.

### Materials and methods

#### Microorganisms and reagents

The microorganisms used in this study are listed in Table 1. All strains were stored frozen at -80 °C in a final concentration of 40% glycerol. L. plantarum FST 1.7 was grown on De Man Rogosa Sharpe (MRS) agar supplemented with 0.05 g/L bromocresol green at 30 °C for 48 h in a micro-aerophilic environment, while S. cerevisiae WLP001, L. fermentati KBI 12.1, and C. subsufficiens C6.1 were grown on yeast potato dextrose (YPD) agar at 25 °C for 48 – 72 h. All microorganisms were stored at 4 °C and sub-cultured

| Table 1 Species, strain designation, and origin of microorganisms used in this study |
|----------------------------------------|-----------------|-----------------|-----------------|
| **Species**                           | **Strain**      | **Origin**      | **References**  |
| Lactiplantibacillus plantarum         | FST 1.7         | UCC Culture Collection (Malted barley) | [26] |
| Lachancea fermentati                  | KBI 12.1        | UCC Culture Collection (Kombucha, USA) | [15] |
| Cyberlindnera subsufficiens          | C6.1            | UCC Culture Collection (Coconut) | [14] |
| Saccharomyces cerevisiae              | WLP001          | California Ale Yeast ®, Whitelabs, San Diego, CA, USA | – |

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weekly. All reagents used in the study were at least analytical grade from Sigma-Aldrich (St Louis, MO, USA) unless stated otherwise.

**Propagation**

Wort was prepared by diluting unhopped Bavarian Pilsner liquid malt extract (Weyermann, Bamberg, Germany) with tap water to a specific gravity of 6 P. The diluted wort was sterilised at 121 °C for 15 min and filtered through sterile Grade IV 320 mm Whatman filter paper (Whatman plc, Maidstone, UK) to remove trub precipitates. *L. plantarum* FST 1.7 was propagated by inoculating a single colony from an MRS agar plate into 10 ml of wort and incubating at 30 °C for 24 h, followed by a 1% sub-culture into 50 ml of fresh wort and incubation for a further 18 h. For propagation of yeast strains, a single colony from a YPD agar plate was transferred into a 250 ml sterile Duran bottle containing 150 ml wort, covered with cotton wool, and incubated for 48 h at 25 °C with an agitation of 100 rpm.

**Fermentation**

Fermentation trials were carried out in sterile 2L Duran bottles filled with 1600 ml of wort and fitted with an airlock. LAB and yeast cells were harvested by centrifugation at 5000 x g for 5 min and washed with sterile tap water to avoid carryover of sugars from the propagation wort. LAB cells were inoculated at a final concentration of 1 x 10⁷ CFU/ml, while yeast cells were pitched at 1 x 10⁶ CFU/ml. Fermentations were performed at 25 °C with each of the three yeast species alone and in combination with *L. plantarum* FST 1.7 (Fig. 1). The NAB was then filled into 330 ml brown FST species alone and in combination with *L. plantarum* fermentations were performed at 25 °C with each of the three yeast strains, a single colony from a YPD agar plate was transferred into a 250 ml sterile Duran bottle containing 150 ml wort, covered with cotton wool, and incubated for 48 h at 25 °C with an agitation of 100 rpm.

**Beer analyses**

At each sampling point, the fermentation bottles were gently shaken to homogenise microbial cells settled at the bottle base into suspension and a 50 ml sample was withdrawn. Cell counts of *L. plantarum* FST 1.7 were determined using MRS agar plates supplemented with 0.05 g/L bromocresol green and 50 µg/ml cycloheximide to suppress yeast growth. Yeast cell counts were determined using YPD agar plates supplemented with 30 µg/ml chloramphenicol to inhibit bacterial growth. pH was measured using a digital pH meter (Mettler Toledo, Columbus, Ohio, USA). Total titratable acidity (TTA) was determined via titration of 10 mL sample against 0.1 M NaOH until pH 7 using an EasyPlus Titratör Easy pH (Mettler Toledo, Columbus, Ohio, USA). Cells were harvested by centrifugation, and the ethanol content (% v/v) and specific gravity (°P) of the supernatant were analysed using density meter DMA 4500 M with Alcolyzer Beer ME (Anton-Paar GmbH, Graz, Austria).

The final beers were analysed as follows. After appropriate dilution and filtration through 0.2 µm syringe-driven filters, organic acids (lactic, acetic, citric, and succinic) were quantified on a Dionex Ultimate 3000 HPLC system (Thermo Fisher Scientific, Waltham, MA, USA), with ultra-violet light/diode array detection (UV/DAD; Thermo Fisher Scientific) and quantification at 210 nm. Analytes were separated on a Hi-Plex H column (8 µm, 7.7 mm × 300 mm; Agilent Technologies, Santa Clara CA, USA) with isocratic elution with 5 mM sulfuric acid and a flow rate of 0.5 mL/min at 60 °C. The compounds were quantified using an external calibration (0.03–6 g/L). After dilution and filtration of the beer samples, sucrose, maltose, glucose, and fructose were quantified via high-performance anion exchange chromatography coupled with pulsed amperometric detection (HPAEC-PAD) on a Dionex ICS-5000+ system (Sunnyvale, CA, USA) equipped with a Dionex CarboPac PA200 column (3 mm × 250 mm) and the corresponding guard column (3 mm × 50 mm; Thermo Fisher Scientific, Waltham, MA, USA) as described by Ispiryan et al. [27]. External standard calibrations were used for each of the compounds between 0.05–1 mg/L and 1–20 mg/L. A 50 mg/L sodium azide solution was used for all sample and reference standard dilutions.

Free vicinal diketones were analysed by MS-Omics. Briefly, samples were mixed with methyl tert-butyl ether (MTBE) 1:1 and in GC–MS vials mixed thoroughly for 2 min, followed by centrifugation for 2 min. All samples were analysed in a randomized order. Analysis was performed as described by Pinu and Villas-Boas [28] with an Agilent 7890B gas chromatograph coupled to an Agilent 5977B quadrupole detector (Agilent Technologies, CA, USA). The system was controlled by ChemStation (Agilent Technologies, CA, USA). Raw data were converted to netCDF format using ChemStation, before the data were imported and processed in Matlab R2014b (Mathworks, Inc.) using tailormade in-house scripts and the PARADISe software described by Johnsen et al. [29]. Free amino nitrogen (FAN) was measured according to MEBAK 2.6.4.1, a ninhydrin-based photometric method where absorbance is measured at 570 nm against glycine. Glycerol was determined via a glucokinase enzymatic assay using a commercial kit (K-GCROLGK, Megazyme International, Wicklow, Ireland).

**Sensory evaluation**

An experienced sensory panel of 8 individuals (5 female and 3 male, age: 23–33) collected descriptors that characterised the aroma and flavour of the NABs. Following this, the
panel evaluated the intensity of the flavour attributes ‘sour’, ‘honey’, and ‘fruity’ and the aroma attributes ‘sweet’ and ‘fruity’ on a scale from 0, ‘not present’ to 10, ‘extremely’. 50 ml samples were provided at ambient temperature (20 °C) in clear glasses labelled with a three-digit code. Sensory evaluations were performed in duplicate under white light and evaluators were instructed to use still water provided to cleanse their palates between samples.

**Statistical analysis**

Fermentations and analyses were performed in triplicate, unless stated otherwise. Values are presented as means ± standard deviation. Statistical analysis was carried out using the Statistical Package for the Social Sciences (SPSS v2.5). Data were analysed using one-way analysis of variance (ANOVA) followed by Tukey’s post hoc test for multiple comparisons. Where equal variances were not assumed, data were analysed by Welch’s ANOVA followed by Games–Howell for pairwise comparison. The level of statistical significance for all tests was set at $p \leq 0.05$.

**Results**

**Fermentation kinetics**

In this study, wort was fermented with *S. cerevisiae* WLP001, *L. fermentati* KBI 12.1, and *C. subsufficiens* C6.1 yeasts, singly and in combination with *L. plantarum* FST 1.7 to produce non-alcoholic beer (NAB). Pre-trials were performed to determine the appropriate fermentation cut-off points to ensure that the ethanol level did not exceed the threshold of 0.5%. Fermentations with *S. cerevisiae* WLP001 and *L. fermentati* KBI 12.1 yeast were stopped after 17 h and 24 h, respectively. As *C. subsufficiens* C6.1 produced low amounts of ethanol which did not exceed the ethanol threshold, fermentations with this yeast strain were stopped after 96 h, prior to the development of an unpleasant solvent-like flavour.

The characterisation of the fermentation wort is shown in Table 2. pH, TTA, alcohol, and extract values were measured continuously at time intervals throughout the fermentations (Fig. 1). In the case of WLP001 and WLP001 + FST 1.7, no change in pH and TTA was observed during the first 8 h of fermentation. After 14 h of fermentation, the pH of WLP001 had decreased to 4.60, while a significantly lower pH of 3.89 was observed for WLP001 + FST 1.7. Only minor changes in pH were observed during the remaining fermentation period, with final values of 4.43 and 3.79 determined for WLP001 and WLP001 + FST 1.7, respectively, after 17 h. The TTA values of WLP001 and WLP001 + FST 1.7 were comparable for the first 14 h of fermentation, after which differences in acidity became evident, with the final TTA of WLP001 + FST 1.7 (2.19 ml) being significantly higher than that of WLP001 (1.26 ml). Drop in extract was comparable for both conditions, with a decrease of approx. 0.81°P observed at the end of fermentation. Similarly, no difference in alcohol production between WLP001 and WLP001 + FST 1.7 was observed, with both fermentations reaching an ethanol level of 0.47–0.49% prior to cut-off.

Similar trends were observed with KBI 12.1 and KBI 12.1 + FST 1.7, with little change in pH or TTA observed during the first 8 h of fermentation. Subsequently, a sharp decrease in pH was observed in both conditions, with KBI 12.1 + FST 1.7 expectedly reaching a lower pH (3.54) than KBI 12.1 (3.89) at the end of fermentation. A corresponding increase in TTA was observed, with NaOH volumes of 2.96 and 3.65 ml required for KBI 12.1 and KBI 12.1 + FST 1.7, respectively. Extract decreased in a linearly in both fermentations, with only minor differences observed in the final extract values for KBI 12.1 (5.43°P) and KBI 12.1 + FST 1.7 (5.38°P). Due to the lower rate of ethanol production by KBI 12.1, the fermentations proceeded for 24 h compared to 17 h for WLP001 fermentations. No significant differences in ethanol production were observed between both fermentations, with final values of 0.42% (KBI 12.1) and 0.45% (KBI 12.1 + FST 1.7) determined.

Despite the longest fermentation time, the highest pH value was observed with C6.1, decreasing to just 4.60 after 96 h, while a final TTA value of 1.47 ml was recorded. In contrast, co-fermentation of C6.1 with FST 1.7 resulted in the lowest pH (3.21) and highest TTA (3.77 ml) values of any of the fermentations performed. Extract values of C6.1 and C6.1 + FST 1.7 decreased by just 0.28°P and 0.38°P, respectively, during fermentation with final values of 5.76°P and 5.61°P measured after 96 h. At the end of fermentation,
C6.1 had produced 0.26% ethanol, while a slightly higher value of 0.32% was recorded for C6.1 + FST 1.7.

Microbial cell growth during fermentations is shown in Fig. 2. Yeasts were pitched at 1 × 10^6 CFU/ml, while L. plantarum FST 1.7 was inoculated in co-fermentations at a concentration of 1 × 10^7 CFU/ml. Results showed that regardless of whether the selected yeasts were fermented singly or in co-fermentation with L. plantarum FST 1.7, significant differences in yeast cell numbers were not observed. Cell counts of WLP001 increased by approx. 1 log CFU/ml after 14 h of fermentation and remained relatively stable thereafter, with final cell numbers of 6.93 log CFU/ml and 6.83 log CFU/ml recorded for S. cerevisiae WLP001 alone and in co-fermentation, respectively (Fig. 2A). A similar trend was observed for KBI 12.1, whereby cell counts increased by 1 log CFU/ml, reaching peak numbers of 7.12–7.16 log CFU/ml after 20 h fermentation and decreasing slightly thereafter (Fig. 2B). Growth of C. subsufficiens C6.1 was slower in comparison to the other yeasts, with increases of 0.5 log CFU/ml and 0.8 log CFU/ml recorded after 48 h and 96 h fermentation, respectively (Fig. 2C) Growth of L. plantarum FST 1.7 was comparable in all co-fermentations, increasing by approx. 1 log CFU/ml with peak cell numbers of 8.10–8.25 log CFU/ml determined.

Analysis of final beers

Sugars and organic acids

The attributes of the final beers are shown in Table 3. WLP001 and WLP001 + FST 1.7 showed similar sugar consumption patterns, with sucrose and fructose completely depleted in both cases, and low levels of fructose remaining (0.54–0.64 g/L). While sucrose was completely depleted during fermentation with L. fermentati KBI 12.1, residual amounts of glucose (0.41 g/L) were detected at the end of the 24 h fermentation period. In contrast, both sucrose and glucose were fully depleted when L. fermentati KBI 12.1 was co-fermented with L. plantarum FST 1.7. Data showed that sucrose was fully depleted after fermentation with C6.1, with high levels of residual glucose (2.47 g/L) detected after 96 h fermentation. In comparison, co-fermentation of C6.1 with L. plantarum FST 1.7 resulted in increased glucose consumption, with a lower level of 1.52 g/L measured in the final beer. Fructose was not fully depleted in any of the fermentations, with levels ranging from 0.5 to 0.9 g/L in the beer samples. The highest levels of maltose depletion were visible in the WLP001 (21.41 g/L) and WLP001 + FST 1.7 (20.14 g/L) fermentations, with approx. 3.15 g/L and 4.42 g/L of this sugar metabolised, respectively. In contrast, residual levels of maltose in the non-S. cerevisiae yeast fermentations were within the range of 22–24 g/L. With regards to maltotriose, levels in the final beers were comparable across all samples, with values of 6.2–7.2 g/L detected. Lactic acid was not detected in the NAB beers fermented with S. cerevisiae WLP001 or C. subsufficiens C6.1 alone, while 0.41 g/L of lactic acid was measured in the L. fermentati KBI 12.1 NAB. As expected, the NABs produced as a result of yeast and L. plantarum FST 1.7 co-fermentation had significantly higher levels of lactic acid present than those produced by yeast fermentation alone. Fermentation of WLP001 + FST 1.7 for 17 h resulted in the production of 0.62 g/L lactic acid, while 1.13 g/L of lactic acid was produced after 24 h fermentation of KBI 12.1 + FST 1.7. The highest level of lactic acid was produced by co-fermentation of C6.1 + FST 1.7, with 2.94 g/L present in the final beer. Citric acid present in the initial wort (0.11 g/L) was not utilised by S. cerevisiae WLP001 and was depleted in varying amounts by WLP001 + FST1.7, KBI 12.1, and KBI 12.1 + FST 1.7, while the acid was fully metabolised by C. subsufficiens C6.1 and C6.1 + FST 1.7. Similarly, more than half (57%) of the acetate acid naturally present in the starting wort was depleted at the end of fermentation with C6.1 and C6.1 + FST 1.7.

Nitrogen metabolism and glycerol

S. cerevisiae WLP001 utilised approx. 25% of the FAN content of the fermentation wort, while WLP001 + FST 1.7 had a slightly higher consumption level of approx. 27%. A similar trend was observed for KBI 12.1 and KBI 12.1 + FST 1.7 fermentations whereby around 22% and 31% of the initial FAN content was depleted by the end of each respective fermentation. The FAN content of the C. subsufficiens C6.1 NAB (83.42 mg/L) was significantly higher than that of the other samples, with just over 3% of the initial amount utilised during the 96 h fermentation period; FAN consumption increased to approx. 25% when C. subsufficiens C6.1 was co-fermented with FST 1.7. Fermentations with S. cerevisiae WLP001 and L. fermentati KBI 12.1 yeasts alone resulted in the production of 0.50 g/L and 0.52 g/L glycerol, respectively, while lower levels of glycerol were present when the yeasts were co-fermented with L. plantarum FST 1.7 (0.40 g/L and 0.42 g/L, respectively). Beers fermented with C6.1 and C6.1 + FST 1.7 had significantly lower levels of glycerol than the other samples, with values of 0.16 and 0.14 g/L determined in the final beers, respectively.

Volatile compounds

The final beers were analysed for fermentation metabolites (Table 4). In terms of higher alcohols, isobutanol concentrations ranged from 1.07 g/L (KBI 12.1) to 1.41 g/L (C6.1 + FST 1.7). C6.1 and C6.1 + FST 1.7 produced significantly lower concentrations of 2-methyl-1-butanol at 0.68 and 0.79 g/L, respectively, than the other samples (1.04–1.22 g/L). Both WLP001 and WLP001 + FST 1.7
produced approx. 9 mg/L of isoamyl alcohol, significantly more than that of the fermentations with non-Saccharomyces yeasts (2.2–6.4 mg/L). The highest concentrations of 2-phenyl ethanol were produced by KBI 12.1 (3.50 mg/L) and KBI 12.1 + FST 1.7 (5.21 mg/L), values almost sevenfold and tenfold higher, respectively, than the concentrations determined in beers produced using C. subsufficiens C6.1 yeast. Levels of total higher alcohols in beers fermented with S. cerevisiae WLP001 and L. fermentati KBI 12.1 yeasts (alone and in co-fermentation with L. plantarum FST 1.7) ranged from 10.40 to 13.96 mg/L, significantly higher concentrations than detected in C6.1 (4.60 mg/L) and C6.1 + FST 1.7 (5.18 mg/L) beers. Total levels of higher alcohols were higher in the beers co-fermented with L. plantarum FST 1.7; however, this difference was not statistically significant. Regarding ester production, C6.1 + FST 1.7 produced
relatively high amounts of ethyl acetate at 19.42 mg/L, almost twofold more than C6.1 alone (9.98 mg/L) and 50–80 times more than WLP001 (0.24 mg/L) and KBI 12.1 + FST 1.7 (0.22 mg/L) yeasts singly or with L. plantarum FST 1.7. Similarly, levels of isoamyl acetate produced by C6.1 and C6.1 + FST 1.7 were significantly higher than in the other beers. Ethyl hexanoate was present in small amounts (0.06 mg/L) in beers fermented with WLP001 and WLP001 + FST 1.7 and was determined in the other beers, but was present in levels below the limit of detection of 0.05 mg/L. Ethyl octanoate was detected in all samples in small amounts, with concentrations ranging from 0.22–0.36 mg/L. Levels of 2-phenylethyl acetate were below the limit of detection in beer produced by S. cerevisiae WLP001 alone, while concentrations of the compound were below the limit of detection in beer produced by C. subsufficiens C6.1 alone. Co-fermentation of S. cerevisiae WLP001 with L. plantarum FST 1.7 resulted in an increase in acetoin levels from 2.36 to 14.41 mg/L, while a twofold increase from 2.69 to 6.19 mg/L was observed in the case of L. fermentati KBI 12.1 co-fermentation. In contrast, acetoin concentrations did not differ significantly between C6.1 and C6.1 + FST 1.7 NABs. Acetaldehyde contents in all beers were well below the flavour threshold of 10 mg/L, with concentrations ranging from 2.26 mg/L (C6.1) to 3.27 mg/L (KBI 12.1).

**Sensory evaluation**

For evaluation of the NABs, a sensory trial was conducted with 8 trained panelists, whereby they were first asked to describe the samples in their own words, followed by an
respective sensory thresholds are highlighted in bold. LOD: limit of detection. Sensory threshold values were obtained from Michel et al. [10]. Values which are above their sensory perception.

FST 1.7 resulted in the beer being perceived as sourer compared to C6.1 (6.6 out of 10). C6.1 + FST 1.7 had a slightly lower acceptability at 6.0 out of 10, with several panelists commenting that the beer was ‘too sour’. WLP001 + FST 1.7 and KBI 12.1 + FST 1.7 NABs produced as a result of co-fermentation were evaluated as being sourer than their WLP001 and KBI 12.1 counterparts, however, not significantly. With regards to beers produced with C. subsufficiens yeast, 66% and 75% of panelists described C6.1 NAB as having a ‘fruity’ aroma and flavour, with notes of pineapple, pear and melon perceived. Similar ‘fruity’ and ‘citrus’ flavours and aromas were described for C6.1 + FST 1.7 beer. Correspondingly, the average intensities of a ‘fruity’ flavour (6.8–7.0 out of 10) and ‘fruity’ aroma (6.5–7.3 out of 10) in both C6.1 and C6.1 + FST 1.7 beers were higher than that of all the other beers (3.1–5.3 out of 10), significantly so (p < 0.05) in the case of WLP001 and WLP001 + FST 1.7. Over 75% of the panelists described C6.1 + FST 1.7 as ‘sour’, with an average intensity of 8.4 out of 10 recorded for this attribute, significantly higher than the other beers (p < 0.01). Consequently, this beer was also perceived as significantly less ‘sweet’ tasting than the other samples (p < 0.05). Regarding overall acceptability, KBI 12.1 + FST 1.7 was ranked the highest with an average rating of 7.0 out of 10, followed closely by L. fermentati KBI 12.1 (6.8 out of 10) and C. subsufficiens C6.1 (6.6 out of 10). C6.1 + FST 1.7 had a slightly lower acceptability at 6.0 out of 10, with several panelists commenting that the beer was ‘too sour’. WLP001 and WLP001 + FST 1.7 had comparable acceptability scores of 6.8 and 6.0 out of 10, respectively. However, it should be

### Table 4: Analysis of volatile compounds in final beers

| Metabolite          | WLP001 S. cerevisiae | WLP001 + FST 1.7 S. cerevisiae + L. plantarum | KBI 12.1 L. fermentati + L. plantarum | KBI 12.1 + FST 1.7 L. fermentati + L. plantarum | C6.1 C. subsufficiens | C6.1 + FST 1.7 C. subsufficiens + L. plantarum | Sensory threshold (mg/L) |
|---------------------|----------------------|-----------------------------------------------|--------------------------------------|------------------------------------------------|-----------------------|-----------------------------------------------|--------------------------|
| Isobutanol          | 1.37 ± 0.13abc       | 1.31 ± 0.11ab                                 | 1.07 ± 0.07a                         | 1.21 ± 0.15ab                                  | 1.21 ± 0.15ab         | 1.41 ± 0.08b                                   | 100                      |
| 2-Methyl-1-butanol  | 1.14 ± 0.13bc        | 1.22 ± 0.15c                                  | 1.04 ± 0.11abc                       | 1.20 ± 0.16c                                   | 0.68 ± 0.18a          | 0.79 ± 0.07bc                                   | 50–70                    |
| Isoamyl alcohol     | 9.05 ± 0.76c         | 9.03 ± 0.67c                                  | 4.80 ± 0.54b                         | 6.36 ± 0.74b                                   | 2.24 ± 0.58a          | 2.54 ± 0.25a                                   | 50–65                    |
| 2-Phenyl ethanol    | 1.37 ± 0.24abc       | 1.65 ± 0.14b                                  | 3.50 ± 0.44d                         | 5.21 ± 0.58d                                   | 0.48 ± 0.12a          | 0.45 ± 0.05a                                    | 125                      |
| Σ Higher alcohols   | 12.93 ± 1.27abc      | 13.20 ± 1.05b                                 | 10.40 ± 1.15c                        | 13.96 ± 1.61c                                  | 4.60 ± 0.98a          | 5.18 ± 0.46a                                    | –                       |
| Ethyl acetate       | 0.24 ± 0.01a         | 0.22 ± 0.03a                                  | < LOD                                | 0.35 ± 0.05a                                   | 9.98 ± 3.79b          | 19.42 ± 2.71c                                  | 33                       |
| Isoamyl acetate     | 0.16 ± 0.06a         | 0.19 ± 0.08a                                  | 0.21 ± 0.06a                         | 0.28 ± 0.08a                                   | 0.54 ± 0.13b          | 0.69 ± 0.08b                                    | 1.6                      |
| Ethyl hexanoate     | 0.06 ± 0.00a         | 0.06 ± 0.00a                                  | < LOD                                | < LOD                                           | < LOD                 | < LOD                                          | 0.23                     |
| Ethyl octanoate     | 0.22 ± 0.00b         | 0.29 ± 0.01b                                  | 0.25 ± 0.00c                         | 0.30 ± 0.01b                                   | 0.33 ± 0.02bd         | 0.36 ± 0.00d                                    | 0.90                     |
| 2-Phenylethyl acetate| < LOD                | 0.13 ± 0.00a                                  | 0.13 ± 0.00a                         | 0.14 ± 0.00a                                   | 0.18 ± 0.02ab         | 0.18 ± 0.01b                                    | 3.8                      |
| Σ Esters            | 0.67 ± 0.03a         | 0.75 ± 0.10a                                  | 0.96 ± 0.09a                         | 0.93 ± 0.14a                                   | 10.88 ± 0.96b         | 20.47 ± 2.79a                                   | –                       |
| Diacetyl            | 0.35 ± 0.01a         | 0.29 ± 0.01b                                  | 0.14 ± 0.03e                         | 0.03 ± 0.00b                                   | < LOD                 | 0.02 ± 0.00d                                    | 1.0–1.15                 |
| Acetoin             | 2.36 ± 0.22a         | 14.41 ± 0.23c                                 | 2.69 ± 0.22a                         | 6.19 ± 0.43b                                   | 3.16 ± 0.60a          | 2.87 ± 0.15a                                    | 50                       |
| Acetaldehyde        | 2.97 ± 0.00a         | 2.38 ± 0.10a                                  | 3.27 ± 0.45a                         | 2.67 ± 0.18a                                   | 2.26 ± 0.54a          | 2.77 ± 0.24a                                    | 10–25                    |

Values are given as means ± standard deviation. Different letters in superscript within the same row indicate a significant difference between the samples (p < 0.05). LOD: limit of detection. Sensory threshold values were obtained from Michel et al. [10]. Values which are above their respective sensory thresholds are highlighted in bold.
noted that differences in overall acceptability scores were not statistically significant.

**Discussion**

Limited fermentation and the use of non-*Saccharomyces* yeasts are techniques that can be used for the production of NAB; however, such processes often result in a sweet-tasting beer with off-flavours, and which is susceptible to microbial spoilage. One approach to address this is the co-fermentation of yeast with LAB during the beer production process. To date, research on the production of beer by co-fermentation of yeast and LAB is limited, with most studies in the literature relating to sour beer production. For example, Dysvik et al. [19] produced sour beer by pre-acidification of wort with *L. buchneri* for 24 h and subsequent fermentation with *S. cerevisiae*, resulting in sour beer with ethanol contents of 3–4%. The same authors also produced alcoholic sour beer by co-fermentation of *L. brevis, L. plantarum*, or *L. buchneri* with *S. cerevisiae* [21]. Others have also exploited the acid-producing nature of LAB for the production of sour beer [24, 25, 30–32]; however, to our knowledge, co-fermentation of LAB and yeast for the production of NAB has not yet been investigated. The aim of the current study was not to produce a sour beer, but to investigate if lactic acid and other metabolite production by LAB could potentially negate the unfavourable effects of limited fermentation with non-*Saccharomyces* yeasts, thus producing an NAB with improved analytical and sensory attributes.

Fig. 3  Spider web diagram of the means of the descriptive attributes from the sensory evaluation of the final beers. Different letters within the same attribute indicate significant differences ($p < 0.05$)
level for NAB in the current study. As expected, the pH values of the NABs in which yeasts were co-fermented with *L. plantarum* FST 1.7 were lower than the NABs produced with the single yeasts. The pH value of the NAB produced by *L. fermentati* KBI 12.1 was significantly lower than that of the NABs produced by *S. cerevisiae* WLP001 or *C. subsufficiens* C6.1 yeasts, due to the unusual lactic acid-producing nature of the *Lachancea* genus [33]. To date, the factors influencing lactic acid production by *L. fermentati* are poorly understood. A study by Bellut et al. [34] investigated the suitability of alternative substrates for reduced-alcohol beer production, with results showing that lactic acid production by *L. fermentati* KBI 12.1 differed significantly between the various substrates, even with the same fermentation parameters (temperature, time, and substrate) and similar sugar composition and amino acid profiles. Osburn et al. [35] reported a lactate production of 0.90 g/L by a *L. fermentati* strain after 1 month of fermentation in 11.4 °P wort, while Domizio et al. [36] determined 0.24 g/L lactic acid in beer after 10 day fermentation with a *L. thermotolerans* strain. In contrast, another study utilising *L. thermotolerans* for beer production reported fourfold less lactic acid produced (0.06 g/L) after 14 day fermentation [37]. Despite the differing fermentation parameters in each study which are known to significantly affect lactic acid production (pitching rate, temperature, and substrate), the production of 0.41 g/L lactic acid by *L. fermentati* KBI 12.1 in the current study after just 24 h fermentation is remarkable. Interestingly, co-fermentation of *L. fermentati* KBI 12.1 with *L. plantarum* FST 1.7 that resulted in a lactic acid concentration similar to that obtained by Bellut et al. [15] following fermentation of *L. fermentati* KBI 12.1 alone for 7 days, highlighting the efficacy of LAB-yeast co-fermentation for lactic acid production. Notably, the concentration of lactic acid in the C6.1 + FST 1.7 NAB was within the range of that typically reported for alcoholic sour beers in the literature (2–9 g/L) [4, 24, 25, 38, 39]. Indeed, according to Tonsmeire [40], any beer with a pH < 3.9 is considered a sour beer; thus, all the NABs in the current study with the exception of *S. cerevisiae* WLP001 and *C. subsufficiens* C6.1 fermented samples would be classed as such. Although not the purpose of the current study, limited co-fermentation of non-*Saccharomyces* yeasts with *L. plantarum* FST 1.7, or other LAB, could be a promising strategy for the production of NAB sour beer, a research area that remains underexplored.

Reduced pH and the presence of lactic acid did not appear to impact the fermentation kinetics of the yeasts, with no significant differences observed in extract reduction, alcohol production or microbial growth between single fermentation and co-fermentations. Similar observations have been made by Chan et al. [25] and Carvalho et al. [41]; however, this is believed to be highly strain-dependent [42]. Dysvik et al. [19] investigated a range of LAB pre-fermentation methods and reported that the growth kinetics of *S. cerevisiae* were impaired only when the strain was inoculated into beer containing viable *L. buchneri* cells. This in turn lead the authors to believe that it was not the pH, lactic acid concentration or the presence of LAB which negatively impacted yeast growth, but rather the viability of the LAB at the time of yeast inoculation [19]. The production of antifungal compounds by the LAB should also be considered as a possible reason for yeast growth impairment in the study [43, 44].

The yeasts used in this study had previously been characterised as relatively acid-tolerant, with no growth impairment reported at pH values as low as 3.5 for *S. cerevisiae* WLP001 and *L. fermentati* KBI 12.1 and pH 4 for *C. subsufficiens* C6.1 [14, 33]. Microbial growth of *L. plantarum* FST 1.7 also remained stable throughout the co-fermentations. Previous work by Peyer et al. [22] established the ability of *L. plantarum* FST 1.7 to grow well in barley malt extract, reaching the highest cell numbers and lowest pH of the 4 LAB species tested. *L. plantarum* FST 1.7 was also able to better withstand lower pH levels with growth continuing until pH 3.0 was reached, while the other species were inhibited at pH 3.5–4.0, highlighting the suitability of this strain for LAB-yeast co-fermentation to produce NAB.

Sugar consumption by yeast and LAB cultures can majorly influence the sensory attributes of the beer, as fructose is perceived as twofold and threefold sweeter tasting than glucose and maltose, respectively. Thus, not only is the perceived sweetness of the beer influenced by the total level of residual sugars, but also by the ratio in which those sugars the present. In this study, maltotriose was not utilised by the non-*Saccharomyces* yeasts and maltose utilisation was negligible, resulting in slower ethanol production and higher extract values in the NABs. The previously reported glucophilic nature of the yeast strains and *L. plantarum* FST 1.7 were observed [14, 15, 22, 33], with glucose being the preferred sugar source in each fermentation. Interestingly, glucose was fully depleted in the WLP001 + FST 1.7 and KBI 12.1 + FST 1.7 NABs after 17 and 24 h fermentation; however, over 30% of the initial glucose content remained in the C6.1 + FST 1.7 NAB after 96 h fermentation. Fructose levels remained relatively unchanged compared to initial wort levels, with no significant consumption of the sugar observed during any of the fermentations, even when glucose was fully depleted. This is in contrast to observations by Bellut et al. [15], whereby fructose was fully depleted along with glucose after 24 h fermentation with *S. cerevisiae* WLP001 or *L. fermentati* KBI 12.1, although it should be noted that the limit of quantification was much higher at 90 mg/L compared to 0.5 mg/L in the current study. In another study, the authors reported that fructose levels after 24 h fermentation with *L. fermentati* KBI 12.1 were similar to the starting concentration in the wort, and fully depleted after 36 h fermentation; hence, it is likely that the short fermentation time in the current study...
interrupted assimilation of the sugar. Additionally, it has been shown that substrate utilisation by yeast is highly dependent on fermentation parameters, such as temperature, pitch rate, wort extract, and so on [14, 33]. In the case of C6.1 and C6.1 + FST 1.7 NABs, fructose was not utilised due to the continued presence of glucose until the end of the fermentation; similar trends have been observed in other studies of Cyberlindnera spp. fermentation [14, 45]. It should also be considered that fermentation is highly depended on the sugar composition of wort, which can be altered to achieve different levels of fermentable and unfermentable sugars. Endres et al. [46] recently developed a single-step mashing procedure at 72 ºC, a temperature at which β-amylase is denatured, but the heat-stable enzyme α-amylase retains its activity, resulting in a wort with decreased glucose and maltose contents and increased levels of the unfermentable sugars maltotetraose and maltotetraose. Using this, a reduced fermentation time of 6 days and a final ethanol concentration of 3.4% were obtained in comparison to a 10–11 day fermentation and an ethanol content of 4.4% with conventional wort. Bellut et al. [34] observed limited ethanol production during fermentation of wort produced from alternative substrates (pulses, cereals, and pseudocereals), due to the low levels of fermentable sugars. Such strategies could be promising for the production of NAB by limited co-fermentation of LAB and yeast, allowing for longer fermentation times and decreased ethanol production.

Free amino nitrogen (FAN) is a measure of the nitrogenous compounds in beer, comprising amino acids, ammonia, small peptides, and individual alpha-acids. A minimum FAN content of 100–130 mg/L in fermentation wort is recommended, while 200–250 mg/L regarded as the optimum level, with insufficient amounts resulting in decreased fermentation rates and limited attenuation [47, 48]. Although the FAN content of the wort in the current study was lower than the recommended value, it is clear from high residual FAN contents of the NABs that the nitrogenous compounds were present in sufficient amounts for the limited time the fermentations were allowed to proceed. FAN content did not differ significantly between NABs fermented with S. cerevisiae WLP001 or L. fermentati KBI 12.1, regardless of whether L. plantarum FST 1.7 was present. The lowest level of FAN assimilation was evident in the C. subsufficiens C6.1 NAB with just 3% of the initial amount consumed during the 96 h fermentation. A similar scenario was observed by Bellut et al. [14] with just 10% of wort FAN consumed after 13 days of fermentation with the same strain. In the current study, FAN consumption increased to 25% when C. subsufficiens C6.1 was co-fermented with L. plantarum FST 1.7 indicating that most of the FAN utilised during this fermentation could be attributed to LAB metabolism. L. plantarum has previously been shown to consume between 15 and 32% FAN when fermented for 24–120 h, with consumption highly dependent on the substrate [22, 23, 49]. FAN consumption by L. fermentati KBI 12.1 has been reported to be relatively low at around 20–30% utilisation after 7–13 day fermentation [15, 33]. Approx. 22% of FAN was consumed by the strain in the current study after 24 h, indicating that the majority of nitrogen metabolism may occur during the early stages of the fermentation.

Higher alcohol concentrations in the WLP001 and KB1 12.1 NABs were similar, with slight increases observed when co-fermentation with L. plantarum FST 1.7 occurred; however, this increase was not statistically significant. In contrast, C6.1 and C6.1 + FST 1.7 demonstrated less than half the amount of higher alcohols, mainly due to lower production of isoamyl alcohol (banana, alcoholic flavour) and 2-phenyl ethanol (rose flavour). The trends in isoamyl alcohol production by the yeasts are concurrent with the literature, with S. cerevisiae WLP001 previously shown to produce more of the compound than L. fermentati KBI 12.1 and C. subsufficiens C6.1 [14, 15, 33]. Low production of 2-phenylethyl alcohol by Cyberlindnera spp. has also been documented [45, 50]. Overall, the levels of higher alcohols in each NAB were well below their respective thresholds. Cyberlindnera spp. have been well characterised in terms of their ester-producing abilities [14, 50, 51] with C. subsufficiens C6.1 producing significantly higher levels than the other yeasts in the current study, mainly ethyl acetate. Ethyl acetate is described as having a fruity flavour but also solvent-like, especially in high concentrations [14]. This solvent-like taste was particularly evident in pre-trials following fermentation with C. subsufficiens C6.1 for > 96 h (data not shown), where ethyl acetate likely reached levels exceeding its flavour threshold of 33 mg/L. Interestingly, the concentration of ethyl acetate in the C6.1 + FST 1.7 NAB was almost double that of the C. subsufficiens C6.1 NAB, indicating that co-fermentation had a significant impact on the production of this ester. Depletion of more than half of the acetic acid present in the fermentation wort by C6.1 and C6.1 + FST1.7 suggests that the high levels of acetate esters in these NABs may have been produced as a result of the esterification of acetate with ethanol by the yeast and/or LAB species [52–54]. Alternatively, it has been hypothesised that suboptimal growth conditions (pH, temperature, and limited substrate) may reduce the flux of acetyl-CoA into the yeast tricarboxylic acid cycle (TCA), resulting in an accumulation of acetyl-CoA which reacts with ethanol to form ethyl acetate [52]. Isoamyl acetate, with a threshold of 1.6 mg/L, is described as having a banana aroma. Concentrations produced by C6.1 and C6.1 + FST 1.7 were also higher than in the other NABs; however, they were lower than values reported in the literature for Cyberlindnera spp. (0.70–7.5 mg/L), likely due to the shorter fermentation period [2, 50, 51]. Similar to the higher alcohols, all esters were below their sensory thresholds. However, it should be considered that the reported thresholds are for...
alcoholic beer. As NAB is substantially lower in volatile compounds and ethanol which have masking activities, it is possible that sensory thresholds for non-alcoholic beverages may be lower [50, 55]. Additionally, synergistic effects between compounds may also result in an impact on flavour, even below the threshold [19]. The concentration of diacetyl, a compound which is regarded as undesirable in beer due to its buttery flavour, was above the threshold of 0.1 mg/L in three NABs. Interestingly, the diacetyl levels in _L. fermentati_ KBI 12.1 NAB exceeded the threshold; however, levels were reduced almost threefold and decreased below the threshold after co-fermentation with _L. plantarum_ FST 1.7. Significantly higher levels of acetoin were determined when _S. cerevisiae_ WLP001 and _L. fermentati_ KBI 12.1 were co-fermented with _L. plantarum_ FST 1.7. Diacetyl, acetoin, and butanediol are by-products of yeast fermentation and can also be produced from LAB citrate metabolism [56–58], an organic compound which was naturally present in the wort and utilised during fermentation. α-Acetolactate is a metabolite produced by both yeast and LAB, which once synthesised is unstable and is decarboxylated to acetoin by α-acetolactate decarboxylase (α-ALD) or can be non-enzymatically decarboxylated to diacetyl (under aerobic conditions). Acetoin may also be produced from diacetyl by diacetyl reductase (DAR) [56–58]. Studies have shown that α-acetolactate synthesis is enhanced at acidic pH [59, 60], a possible explanation as to why acetoin concentrations were increased in the LAB co-fermented beers. After primary fermentation, yeasts have the capability to reabsorb diacetyl and produce 2,3-butanediol via acetoin [47]. From the results, it is clear that this reaction was not complete in the case of WLP001 + FST 1.7 and KBI 12.1 + FST 1.7 NABs; however, the lower levels of both diacetyl and acetoin in the C6.1 + FST 1.7 NAB suggested that the longer fermentation time allowed completion of this conversion.

Although below their respective thresholds, the increased ester production by _C. subsufficiens_ C6.1 was clearly detected in the sensory evaluation, with C6.1 and C6.1 + FST 1.7 NABs ranking highest in fruity taste and aroma, and assigned descriptions, such as ‘pear’, ‘melon’, and ‘pineapple’. Despite having the highest levels of residual sugars, C6.1 NAB was not evaluated as significantly sweeter smelling or tasting than the other samples, indicating that the fruity flavour and aroma masked the wort-like off-flavour. Additionally, the main residual sugar in this sample was maltotriose which is significantly less sweet tasting than the other carbohydrates in the beer. Due to the intense acidity of the C6.1 + FST 1.7 beer, it was ranked as the least sweet tasting; however, the acceptability of this sample was decreased compared to C6.1 NAB, with some panelists considering it ‘too sour’. However, it should be considered that a decreased co-fermentation time of less than 96 h could potentially result in an NAB with a fruity flavour and a balanced ratio of acidity and sweetness, without being perceived as overly sour. Alternatively, the intense sourness of this sample could be exploited to produce a sour NAB, research on which is sparse. Of note is the significant reduction in the perceived honey flavour of the WLP001 and KBI 12.1 beers which were co-fermented with _L. plantarum_ FST 1.7, underlining the previously highlighted contribution of LAB fermentation to the sensory properties of beer [61–63]. KBI 12.1 + FST 1.7 was ranked as the most acceptable beer in the sensory evaluation. The lactic acid content of 1.13 g/L in the NAB far exceeded the sensory threshold of 80 mg/L; this was perceived in the sensory evaluation with this beer being ranked as sourer than the non-co-fermented beer; however, the difference was not statistically significant. Additionally, the increased sourness did not appear to be off-putting to the panelists and did not negatively affect the acceptability of the product.

**Conclusion**

In summary, co-fermentation of _L. fermentati_ KBI 12.1 and _C. subsufficiens_ C6.1 with _L. plantarum_ FST 1.7 was found to be a promising strategy for the production of NAB (<0.5% ethanol). Due to the limited fermentative capacity of the yeasts and subsequently lower ethanol production, fermentation time could be extended beyond that of which was possible with the control yeast, _S. cerevisiae_ WLP001. Short co-fermentation of _L. fermentati_ KBI 12.1 with _L. plantarum_ FST 1.7 produced an NAB with a relatively high concentration of lactic acid, lower levels of residual sugars, and a reduction in the amount of diacetyl to below the sensory threshold. Moreover, the beer was rated the most acceptable in the sensory evaluation. NAB produced by C6.1 + FST 1.7 was described as having a fruity taste and aroma which masked the wort-like flavours/aromas due to the high levels of residual sugars. The high concentration of lactic acid in this sample resulted in it being evaluated as too sour by some panelists; however, this could be exploited to produce an NAB sour beer or alternatively, future work could investigate the alteration of the fermentation parameters to produce a fruity beer with a more balanced acidity. Additionally, further work could explore the impact of different LAB species and strains on the fermentation kinetics and sensory attributes of NABs. Overall, this study highlights the potential of limited co-fermentation of non- _Saccharomyces_ yeasts and LAB for application in non-alcoholic beer brewing.

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Declarations

Conflict of interest The authors declare that there are no conflicts of interest.

Compliance with ethics requirement This article does not contain any studies with human or animal subjects.

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