Long-insert clone experimental evidence for assembly improvement and chimeric chromosomes detection in an allopentaploid beer yeast

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Abstract

Lager beer is made with the hybrid Saccharomyces pastorianus. Many publicly available S. pastorianus genome assemblies are highly fragmented due to the difficulties of assembling hybrid genomes, such as the presence of homeologous chromosomes from both parental types, and translocations between them. To improve the assembly of a previously sequenced lager yeast hybrid Saccharomyces sp. 790 and elucidate its genome structure, we proposed the use of alternative experimental evidence. We determined the phylogenetic position of Saccharomyces sp. 790 and established it as S. pastorianus 790. Then, we obtained from this yeast a bacterial artificial chromosome (BAC) genomic library with its BAC-end sequences (BESs). To analyze these data, we developed a pipeline (applicable to other assemblies) that classifies BES pairs alignments according to their orientation. For the case of S. pastorianus 790, paired-end BESs alignments validated parts of the assembly and unpaired-end ones suggested contig joins or misassemblies. Importantly, the BACs library was preserved and used for verification experiments. Unpaired-end alignments were used to upgrade the previous assembly and provided an improved detection of translocations. With this, we proposed a genome structure of S. pastorianus 790, which was similar to that of other lager yeasts; however, when we estimated chromosome copy number and experimentally measured its genome size, we discovered that one key difference is the outstanding S. pastorianus 790 ploidy level (allopentaploid). Altogether, our results show the value of combining bioinformatic analyses with experimental data such as long-insert clone information to improve a short-read assembly of a hybrid genome.

Keywords: lager; yeast; hybrid; genome; BAC; assembly; translocations

Introduction

There are two main types of beers depending on the kind of yeast used for its production. Ale beer is elaborated with some strains of Saccharomyces cerevisiae, a well-studied organism (Goffeau et al. 1996; Engel et al. 2014), whereas lager beer is produced with Saccharomyces pastorianus (Martini and Kurtzman 1985; Martini and Martini 1987). Most large breweries utilize lager yeast, hence the importance to study this organism for industrial application. S. pastorianus is a hybrid between an S. cerevisiae (Engel et al. 2014; Goffeau et al. 1996) probably of ale origin, and Saccharomyces eubayanus (Baker et al. 2015; Libkind et al. 2011), another closely related wild yeast that has been isolated from multiple locations around the world (Libkind et al. 2011; Bing et al. 2014; Peris et al. 2014; Gayevskiy and Goddard 2016). All of these S. eubayanus isolates group into two principal populations potentially originating from Patagonia, one of which the other globally distributed subpopulations derive (Peris et al. 2014, 2016; Eizaguirre et al. 2018).

Even though the highest number of subpopulations, genetic diversity, and abundance have been found in South America (Eizaguirre et al. 2018; Langdon et al. 2020; Nespolo et al. 2020), based on higher genetic similarities and experimental observations, an Holartic subpopulation seems to be the most likely parent of the lager beer yeast (Bing et al. 2014; Peris et al. 2016; Brouwers et al. 2019). A third type of beer, belonging to traditional Belgian beers, fermented with S. cerevisiae and Saccharomyces kudriavzevii hybrids can be considered (González et al. 2008; Gallone et al. 2019; Langdon et al. 2019). Furthermore, other hybrids, even multiple hybrids, have been isolated from different fermentations, such as wine and cider (Masneuf et al. 1998; González et al. 2006; Peris et al., 2012), evidencing the important role of hybridization for evolution and environment adaption (Gallone et al. 2019; Langdon et al. 2019).

The hybrid nature of S. pastorianus implies that its genetic content can be segregated in two parental sub-genomes. After the
DNA repeats, and simultaneously, assembly algorithms fail in
bleeding into longer contiguous sequences or contigs. Nonetheless,
strains (Hewitt
the genome structure and chromosome copy number of
rental-like genomes or genome sequence references to determine
The initial sequencing results, or reads, have been aligned to pa-
number of copies; otherwise, they cannot be detected by this
requires that the involved chimeric chromosomes have a different
Changes in sequencing depth across the chromosome indicate re-
Reads covering a nucleotide base, or depth, in a chromosome re-
and group 2 or Frohberg, that have different phenotype, geographical
origin, and genetic elements distribution (Liti et al. 2005; Dunn and Sherlock 2008; Gibson et al. 2013). Yeasts from the dis-
tinct lager groups are thought to share one common origin due to the
presence of conserved recombination points across several group-1 and group-2 strains (Hewitt et al. 2014). Although, the
specific origin of each lager group has been highly debated, from
hypotheses suggesting one hybridization and later divergence,
two independent hybridization events, or more complex phyloge-
etic stories (Dunn and Sherlock 2008; Hewitt et al. 2014; Okuno et al. 2016; Monerawela and Bond 2017; Salazar et al. 2019).
Nonetheless, the type and combination of chromosomes, or gen-
ome structure, varies greatly among lager strains, being more
notorious between yeasts from different groups (Monerawela and Bond 2017; Gallone et al. 2019). Similarly, the total DNA content,
or ploidy level, also fluctuates in lager yeasts due to changes in
chromosomes copy number; however, it is generally acknowl-
edged that group-1 yeasts are allotriploids, whereas group-2 are
alloetraploids (Walther et al. 2014; van den Broek et al. 2015). The
overall genome structure and chromosome copy number is thought to be unique for each lager yeast isolate and responsible for the
observed differences in brewing-related traits (van den Broek et al. 2015). Therefore, knowing the genome structure of lager yeasts is important to characterize unstudied or newly iso-
strains, to associate their genotype with their phenotype, and to serve as a background for laboratory experiments design,
yeast breeding, and selection programs.
Massive parallel or short-reads sequencing allowed to study the
genome of several S. pastorianus yeast strains (Nakao et al. 2009; Doštálek et al. 2013; Walther et al. 2014; van den Broek et al. 2015; De León-Medina et al. 2016; Okuno et al. 2016; Langdon et al. 2019).
The initial sequencing results, or reads, have been aligned to pa-
rental-like genomes or genome sequence references to determine
the genome structure and chromosome copy number of S. pastoria-
us strains (Hewitt et al. 2014; Walther et al. 2014; van den Broek et al. 2015), based on the principle that the average number of
reads covering a nucleotide base, or depth, in a chromosome re-
gion is proportional to the number of copies of that region. Changes in sequencing depth across the chromosome indicate re-
combination sites. However, the detection of recombination sites requires that the involved chimeric chromosomes have a different
number of copies; otherwise, they cannot be detected by this
method. Furthermore, this process per se does not retrieve the
chromosome sequence; for that, the reads have also been assem-
bled into longer contiguous sequences or contigs. Nonetheless,
most of the publicly available S. pastorianus assembly reports con-
sist of hundreds, if not thousands, of contigs (NCBI Genome, http://www.ncbi.nlm.nih.gov/genome), and the N50 of the assem-
bles, a measure of the assemblies’ contiguity, is usually smaller
than the average size of a yeast chromosome. The above may be
due to intrinsic limitations of the used sequencing and assembly
technologies, as well as the inherent complexity of assembling hy-
brid genomes (Pryszcz and Gabaldón 2016). In more detail, short-
reads produced by massive parallel sequencing cannot span long
DNA repeats, and simultaneously, assembly algorithms fail in
such repeated regions (Schatz et al. 2010; Alkan et al. 2011).
Moreover, highly similar intraparental sequences from nonrecom-
bined and chimeric chromosomes are assembled into one consen-
sus contig, and sequences flanking recombination points are not
linked, producing fragmented assemblies (Pryszcz and Gabaldón 2016). Such assemblies contain small contigs difficult to scrutinize
and use to design experiments.
One method used to order contigs in S. pastorianus assemblies is to utilize synteny, or whole genome alignments, with the pa-
rental genomes solely (Walther et al. 2014). Nonetheless, ordering contigs using a reference disregard the S. pastorianus heterozygo-
city. Another method used to upgrade lager yeast assemblies is scaffolding (van den Broek et al. 2015; Okuno et al. 2016). This pro-
cess uses specially paired massive parallel reads, or mate pairs,
separated by a fixed DNA length, or insert. Then the scaffolding
orders and orients contigs to produce even longer assembled
sequences called scaffolds. However, the mate pair sequencing li-
ibraries possess short inserts that may not span the aforementioned long repeated regions, are often not publicly available,
and/or still may not produce assemblies at chromosome level.
Most recently, long-reads sequencing technologies, such as
Pacific Bioscience and Oxford Nanopore Technologies, have been
used to sequence lager yeast genomes; but until now, only two assemblies of this kind of data are available in the public reposi-
tories (Liu et al. 2018; Salazar et al. 2019).
Regarding the mentioned technologies limitations, we sought
an alternative approach to validate and improve the available
Saccharomyces sp. 790 genome assembly (De León-Medina et al.
2016), a previously sequenced lager yeast hybrid, and also to elu-
cidate its genome structure. We envisioned that this alternative
could be in the form of experimental data. Bacterial artificial chromosomes (BACs) (Shizuya et al. 1992) are an experimental re-
source consisting of circular and stable cloning vectors that can
contain large DNA inserts (up to 300 kbp). The first bases from
both ends of the BACs inserts, or BAC-end sequences (BESs), can
be explored using universal primers anchored in the vector back-
bone and Sanger sequencing. BACs were frequently used to assist
gene sequencing and assembly (Weber and Myers 1997) and
could be applied to investigate an already sequenced and assem-
bled hybrid genome; although, they have never been used for this
particular purpose. BACs are also an alternative source of infor-
mation, since they are created from the same organism but inde-
pendently from the information used for genome assembly.
Furthermore, the features of these vectors, such as the paired-
end, long-insert, and single allele information, make them attrac-
tive to elucidate the genome structure of a hybrid organism, as
we detail in this study. Therefore, we constructed a BAC library of
Saccharomyces sp. 790 and obtained the BESs. As part of our
efforts, we developed, curated, and made available a pipeline for
analyzing the data. Afterward, we used the generated paired-end
and long-insert information to validate and improve the available
Saccharomyces sp. 790 assembly (De León-Medina et al. 2016),
and using the single allele information, we elucidated the
Saccharomyces sp. 790 genome structure. Finally, we observed a
Saccharomyces sp. 790 genome feature that might differentiate it
from other lager yeasts.

Materials and methods
Yeast strain and sequences files
The brewing yeast strain Saccharomyces sp. 790, specific genome
sequences (scaffolds 17 and 22, File S1), and the short-reads se-
quencing library were obtained from Cervecería Cuauhtémoc
Moctezuma S.A. de C.V. (Monterrey, NL, Mexico). The reference sequence of *S. cerevisiae* S288c genome (Engel et al. 2014) was downloaded from the Saccharomyces Genome Database (https://www.yeastgenome.org). Genome sequence assemblies of *S. eubayanus* FM1318 (Baker et al. 2015), 10 *S. pastorianus* strains (Nakao et al. 2009; Dostálek et al. 2013; Walther et al. 2014; van den Broek et al. 2015; Okuno et al. 2016) (Supplementary Table S1), Saccharomyces sp. 790 (De León-Medina et al. 2016), and *S. cerevisiae* sp. M14 (Liu et al. 2018) were retrieved from the NCBI Genome database (NCBI, Genome http://www.ncbi.nlm.nih.gov/genome). See Supplementary Table S1 for more information about the assemblies.

**Genome annotation and phylogenetic analysis**

We annotated the genome assemblies of *S. pastorianus* strains CBS 1503, CBS 1513, CCY48-91, CBS 1483, CBS 1538, WS 34/70, Saccharomyces sp. M14, and Saccharomyces sp. 790 (Dostálek et al. 2013; Walther et al. 2014; van den Broek et al. 2015; De León-Medina et al. 2016; Okuno et al. 2016; Liu et al. 2018) (Supplementary Table S1). Genome annotation of the yeast assemblies was performed with MAKER2 (v2.31.10) (Holt and Yandell 2011). In brief, gene models were obtained with SNAPP (v2013-11-29) (Korf 2004) and Augustus (v3.2.3) (Stanke et al. 2006) as ab initio gene predictors and with homology to the transcripts and proteins of *S. cerevisiae* S288c (Engel et al. 2014) and *S. eubayanus* FM1318 (Baker et al. 2015). For the functional annotation, we used *S. cerevisiae* S288c (Proteome ID: UP000002311) and *S. eubayanus* FM1318 (Proteome ID: UP000052040) proteomes (UniProt, https://www.uniprot.org/). Proteomes were combined in a single file, which means that both parental sequences are aligned competitively and the best hit of a gene corresponds to the most likely function of the gene, but also the most likely parental type. Orthologous genes’ identification from each sub-genome was carried out using the OrthoMCL algorithm (v1.4) (Li et al. 2003) as implement in get_homologues.pl (v23072018) (Contreras-Moreira and Vinuesa 2013) and with the gene information of *S. cerevisiae* S288c (Engel et al. 2014) and *S. eubayanus* FM1318 (Baker et al. 2015) as reference. This led to identify 316 and 1470 one-to-one orthologous groups for the *S. cerevisiae* and *S. eubayanus* sub-genomes, respectively. For each orthologous group, the nucleotide sequences among the eight lager strains and the respective parental sequences were aligned with MAFFT (v7.453) (Katoh and Standley 2013). A concatenated full-length data set containing all orthologous groups for each sub-genome was used for phylogenetic reconstruction. RAxML (v8.2.12) (Stamatakis, 2014) was used to build the maximum likelihood trees based on the GTRGAMMA model with 1000 rapid bootstraps. Both trees were visualized with ggtree (v4.11) (Yu et al. 2017).

**Library construction, characterization, and end-sequencing**

The BAC genomic library and BESs of Saccharomyces sp. 790 were generated at Clemson University Genomics Institute (CUGI, Clemson, SC, USA). The library was built with HindIII, the pLndig0BAC536 vector, and *Escherichia* coli DH10B. Clones were randomly selected, purified, and characterized with NotI (*n* = 45) (New England Biolabs, Ipswich, MA, USA) (Farrar and Donnison 2007). The insert was visualized in a 1% pulsed field gel electrophoresis agarose-0.5X TBE gel (Molecular Sigma Biology, St. Louis, MO, USA) in a CHEF Mapper XA (Bio-Rad, Hercules, CA, USA) at 6.0 V cm⁻¹, 120°, 5–15 s, lineal, 16 h, and 14°C. The estimated insert size was calculated by interpolation to a DNA ladder. The formula \( W = NI/G \) was used to calculate the genomic coverage (W), where \( N \) is the number of clones, \( I \) is the average insert size, and \( G \) is the genome size (the sum of the parental genomes sizes, i.e., 23.8 Mbp). All the clones were subjected to vector purification and sequencing from both ends with Sanger technology using universal primers T7 and M13.

**BESs processing, alignment pipeline, and other bioinformatic procedures**

We designed a pipeline to analyze BESs information (Supplementary Figure S1; scripts available at https://github.com/Lriego/BES-analysis). Input genome files preparation, BESs processing, and alignment were performed with a custom-made BASH script (bes_analysis.sh) as follows: the raw BESs reads in ABI format were converted to FASTQ with Seqret (vEMBOSS : 6.6.0.0) (Rice et al. 2000). Sequence’s quality was assessed with FastQC (v0.11.5) (available online at http://www.bioinformatics. babraham.ac.uk/projects/fastqc). Sequences were trimmed and filtered with Trimmomatic (v0.36) (Boiger et al. 2014) with the following parameters: HEADROP: 91 LEADING: 20 TRAILING: 20 SLIDINGWINDOW: 4:20 MINLEN: 35. The resulting sequences were converted to FASTA format with FASTX-Toolkit (v0.0.14) (available online at: http://hannonlab.cshl.edu/fastx_toolkit/). Sequences from vectors with no insert (vector positive) and sequences from yeast DNA (yeast positive) were detected by aligning the BESs with the pLndig0BAC536 vector sequence, and the *S. cerevisiae* S288c (Engel et al. 2014), *S. eubayanus* FM1318 (Baker et al. 2015), and Saccharomyces sp. 790 (De Leόn-Medina et al. 2016) assemblies using BLAST (v2.6.0+) (Altschul et al. 1990; Camacho et al. 2009) with default parameters and an e-value of \( 1e^{-6} \). BESs were deposited in the DDBJ/EMBL/GenBank database under the accession number LIBGSS_039438. BESs with no insert were removed from the reported data set to comply with the database requirements, but the complete original data set was described and used in subsequent analyses. BESs nucleotide sequences can be located in the corresponding genome assembly using BLAST (v2.6.0+) (Altschul et al. 1990; Camacho et al. 2009), BLAT (v36x2) (Kent 2002), MEGA-BLAST (v2.6.0+) (Camacho et al. 2009), NUCMER (v3.1) (Kurtz et al. 2004), BOWTIE (v1.2.2) (Langmead et al. 2009), BOWTIE2 (v2.3.4.1) (Langmead and Salzberg 2012), and BWA (v0.7.17) (Li and Durbin 2009). All the alignment tools were tested by aligning the BESs with the Saccharomyces sp. genome assembly (De Leόn-Medina et al. 2016) (Supplementary Table S2). A tool presenting intermediate results with respect to the total of aligned BESs, in this case BOWTIE2 (Langmead and Salzberg 2012), was selected to perform all BESs alignments presented in this work, unless otherwise specified. For any tool used, only one alignment per BES, either the longest or primary alignment, is selected by the BASH script. BESs alignment results were converted into GFF table format and screened with a Python script (screening.py). The Python script takes in the GFF file and detects BESs pairs that align in the same sequence. Then, the BESs alignments were classified according to the orientation between the pairs as paired-end, positive, negative, or opposite (Supplementary Figure S2). If only one BES pair aligns, it is classified as singleton. BESs pairs that align in different contigs are classified as unpaired-end (Supplementary Figure S2) and used for downstream analyses. Unpaired-end BESs alignments were categorized according to their hypothetical insert size (the added distance from each unpaired-end BES alignment site to the end of its aligning sequence). The screening Python script outputs a summary table, and a specially formatted GFF file that includes the BESs pairs classification, the insert size, the pair information...
for unpaired-end alignments, and an alignment type color code to aid the visual inspection of the file in the Integrative Genomics Viewer (v2.4.14) (Thorvaldsdóttir et al. 2013). Two additional custom-made Python scripts (sliding_paired.py and sliding_unpaired.py) were designed and used to detect large regions (>60 kb) not spanned by paired-end BESs alignments and regions enriched with unpaired-end BESs pairs pointing to the same contig (consistent). These scripts scan the formatted BESs alignments GFF file using a sliding window algorithm and output a BED file with the coordinates of the detected regions. A visual summary of the results can be obtained with an R script (graphic_summary.R) from the summary table and the BESs scanning results. The integrated information of BESs alignments, BESs scanning results, and annotation were visualized in the Integrative Genomics Viewer (Thorvaldsdóttir et al. 2013) and figures were generated using the R package Gviz (v1.28.3) (Hahne and Ivanek 2016). Scaffolding of the Saccharomyces sp. 790 assembly (De León-Medina et al. 2016) was performed using SSPACE (v3.0) (Boetzer et al. 2011) and the BOWTIE2 BESs alignments results. To propose a genome structure of Saccharomyces sp. 790, the output scaffolds were ordered and joined into chromosomes using NUCMER (v3.1) (Kurtz et al. 2004) and the S. cerevisiae S288c (Engel et al. 2014) and S. eubayanus FM1318 (Baker et al. 2015) genomes as reference. The best mapped location of each query contig (show-tilling -p option) was chosen as best hit. After this process, BESs were realigned to the resulting sequences, and regions enriched with consistent unpaired-end BESs alignments were detected using our scanning script (sliding_unpaired.py) at each of the following two steps. In the first step, the unpaired-end BESs scanning results were used to detect potential recombination sites in the newly joined chromosomes, then the closest gap toward the unpaired-end BESs alignments direction was selected, and the chromosomes were split in the gap position. In the second step, a final genome structure was generated with Biostrings (v2.52.0) (Pagès et al. 2019) using all of the possible homologous chromosomes joins suggested by the unpaired-end BESs scanning results. The Saccharomyces sp. 790 genome structure in FASTA format was annotated identically as the other lager yeast assemblies and plotted to portray the Saccharomyces sp. 790 genome structure using genoPlotR (v0.8.9) (Guy et al. 2010).

Gaps and homologous translocations amplification

Briefly, primers flanking potential contig joins and homologous translocations were designed using the appropriate sequences (Supplementary Table S3). Library clones spanning regions of interest were selected to be used as template (Supplementary Table S3). BACs were purified (Farrar and Donnison 2007). DNA was amplified by PCR using standard conditions, and the resulting fragments were visualized in an electrophoresis gel.

Flow cytometry analysis to measure total DNA content

DNA content of S. cerevisiae CLA1 (Avendaño et al. 1997), S. eubayanus FM1318 (Baker et al. 2015), S. pastorianus strains CBS 1513 (Walther et al. 2014) and CBS 1483 (van den Broek et al. 2015), and the study yeast was measured. S. cerevisiae CLA1 and S. eubayanus FM1318 cells were grown in casamino acids medium (yeast nitrogen base 1.7 gL⁻¹, (NH₄)₂SO₄ 5 gL⁻¹, casamino acids 6 gL⁻¹, and dextrose 20 gL⁻¹), whereas the lager yeasts cells were grown in YPD media (yeast extract 10 gL⁻¹, peptone 20 gL⁻¹, and dextrose 20 gL⁻¹) to avoid nutritional stress. The Haase and Reed (2002) protocol was used to stain yeasts’ DNA with SYTOX Green (Thermo Fisher Scientific, Waltham, MA, USA). Samples were analyzed in the Molecular Biology Division’s (IPICYT) flow cytometer BD FACSCalibur (Becton Dickinson & Company, Franklin Lakes, NJ, USA). Fluorescence was captured in the FL1-A channel and 5000 events in the assigned window were registered. The fluorescence geometric means of GO-1 and G2 cell cycle phases were calculated in the FlowJo X software (v10) (Becton Dickinson & Company, Franklin Lakes, NJ, USA). A linear regression was employed to determine the DNA content of the Saccharomyces sp. 790 strain, considering the fluorescence geometric means and the reported genome size of the control strains (Engel et al. 2014; Walther et al. 2014; Baker et al. 2015; van den Broek et al. 2015). Fluorescence intensity in FLA-1 channel against the number of events (fluorescence histogram) was plotted using the R packages flowCore (v1.50.0) (Hahne et al. 2009) and ggcyto (v1.12.0) (Van et al. 2018).

Bioinformatic estimation of Saccharomyces sp. 790 chromosome copy number

A Saccharomyces sp. 790 short-reads sequencing library was aligned with the parental-like genomes of S. cerevisiae S288c (Engel et al. 2014) and S. eubayanus FM1318 (Baker et al. 2015) using BOWTIE2 (v2.3.4.1) (Langmead and Salzberg 2012). The output file in SAM format was processed in two separate ways. First, to estimate the haploid distribution, the SAM file was converted to BAM using Samtools (v1.7) (Li et al. 2009) and from BAM to BED using Bedtools (v2.26.0) (Quinlan 2014). Next, the sequencing depth histograms per parental chromosome were obtained using Bedtools genomcvc (v2.26.0) (Quinlan 2014). Second, to calculate the sequencing depth mean across each parental chromosome, sequencing depth per base was calculated using Samtools (v1.7) (Li et al. 2009). Results of the first and second analysis were processed with a bespoke R script. The script establishes the smallest distribution in the chromosome sequencing depth histograms as the haploid distribution and calculates the sequencing depth per base mean in 1000 bp windows. At last, the formula \( C = \log(W/H) \) was used to obtain the window copy number \( C \), where \( W \) is the window depth mean and \( H \) is the haploid distribution. It was established that the logarithm base two equal to zero is one copy. All windows copy number for each chromosome were plotted using the R graphics package (v3.6.3).

Data availability

All lager yeast assemblies used in this study are publicly available (see Supplementary Table S1). BESs generated in this study were deposited in the DDBJ/EMBL/GenBank database under the accession number LIBGSS_039348. Curated code to analyze the LIBGSS_039348 is available at https://github.com/Lriego/BES-analysis. Not curated code and raw data are available upon request. The authors affirm that all data necessary for confirming the conclusions of the article are present within the article, figures, and tables. Supplementary material available at figshare: https://doi.org/10.25387/g3.14156912. Supplementary File S1 contains the potential nonhomologous translocation (scaffolds 17 and 22) in FASTA format. File S2 is the BED file of consistent unpaired-end BESs alignments in S. pastorianus 790. Supplementary File S3 is the BED file of consistent interparental unpaired-end BESs alignments in the combined parental genomes of S. cerevisiae S288c and S. eubayanus FM1318 with at least three supporting clones. File S4 is the BED file of consistent interparental unpaired-end BESs alignments in the combined parental genomes of S. cerevisiae S288c and S. eubayanus FM1318 with at least two supporting clones. The lager yeast strain 790 as well as the BAC library are property of Cervecería Cuauhtémoc Moctezuma S.A. de C.V.
Results

Saccharomyces sp. 790 is a group-2 lager yeast

Saccharomyces sp. 790 was established as a hybrid between S. cerevisiae and S. eubayanus (De León-Medina et al. 2016), nonetheless, its phylogenetic position with respect to other hybrids of this type is unknown. To answer the former question and to provide clarity of the source DNA used for the library construction, we performed a phylogenetic analysis using the orthologous genes from each sub-genome of Saccharomyces sp. 790 strain (De León-Medina et al. 2016) and seven larger beer-related strains (Dostálek et al. 2013; Walther et al. 2014; van den Broek et al. 2015; Okuno et al. 2016; Liu et al. 2018) (Supplementary Table S1). In both S. cerevisiae and S. eubayanus orthologous genes trees, all group-1 S. pastorianus strains (Supplementary Table S1) grouped together, and strains classified as group-2 (Supplementary Table S1) were in close distance with each other (Figure 1; Supplementary Figure S3). The phylogenetic position of Saccharomyces sp. M14 was unknown, but in our phylogenetic reconstruction this yeast always clustered with other group-2 yeasts, such as S. pastorianus CBS 1483. Saccharomyces sp. 790 also clustered with other group-2 S. pastorianus strains (Figure 1; Supplementary Figure S3). Based on these results, we classified Saccharomyces sp. 790 as S. pastorianus 790 of group-2.

A redundant, long-insert, and paired-end experimental and sequence resource for studying the S. pastorianus 790 genome

To generate resources for studying the S. pastorianus 790 genome, we constructed from this yeast a BAC genomic library comprised of 2304 clones. The BAC library was preserved and could be used for verification experiments. The average estimated genomic coverage, or redundancy, was 9.0X, the measured clone insert proportion was 100% (n = 45) (Supplementary Figure S4), and the insert size of the library was 93.4 ± 31.7 kbp (Table 1). The observed large insert size could be a key feature to study the S. pastorianus 790 genome at large scale, since it can span conflicting regions of DNA such as repeated sequences or translocation points. We obtained the BESs of the library (4608 total BESs), we filtered the resulting sequences and 4004 passed the quality parameters. From the passing BESs, 3586 were paired (both pairs) and 418 were single (one pair) (Table 1 and Supplementary Table S4). Furthermore, the average sequence length was 710.44 ± 212.24 bp and the total sequenced bases were 2,844,613 bp (Table 1). This sequence information is the link between the BAC library experimental data and the bioinformatic genome information. To detect sequences from vectors with no insert (vector positive), we aligned the 4004 quality BESs with the vector sequence using BLAST (Altschul et al. 1990; Camacho et al. 2009). Of these, 109 aligned exclusively to the vector, which indicated sequences from BACs with no insert, whereas 3895 did not align (Supplementary Table S4). BESs are available in the DDBJ/EMBL/GenBank database under the accession number LIRGSS_039348. BESs with only vector sequences were excluded for a total of 3895 BESs in the reported data set.

BESs alignment pipeline benchmarking with several S. pastorianus assemblies

To investigate our specific type of data, we designed a pipeline for analyzing BESs information. The scripts and pipeline were curated and published for other potential users (https://github.com/LrieCo/BES-analysis). To test the proper functioning of our

Table 1 Genomic library characterization and end-sequencing summary

| BAC-based genomic library |
|---------------------------|
| Clones                    | 2304 |
| Genomic coverage          | 9.0X |
| Clone insert proportion   | 100% |
| Average estimated insert size | 93.4 ± 31.7 kbp |
| n = 45                    |
| BESs                      |      |
| Total BESs                | 4608 |
| Quality BESs              | 4004 |
| Paired BESs               | 3586 |
| Single BESs               | 418  |
| Average sequence length   | 710.44 ± 212.24 bp |
| Sequenced bases           | 2,844,613 bp |

Figure 1 Phylogenetic relationships of Saccharomyces sp. 790 and seven lager beer-related yeast strains. All yeast assemblies were annotated with MAKER2 pipeline (Holt and Yandell 2011) and the orthologous genes from each sub-genome segregated and used to construct specific parental sub-genome trees. (A) Maximum likelihood tree of the S. cerevisiae parental sub-genome genes. (B) Maximum likelihood tree of the S. eubayanus parental sub-genome genes. In both trees, Saccharomyces sp. 790 clustered closer to other group-2 lager yeast strains; therefore, we classified it as S. pastorianus 790 of group-2. Numbers at the nodes are the percent of bootstrap replicates that support it (only numbers higher than 50% are shown). The scale on the bottom left represents genetic distances in substitutions per nucleotide. Numbers in parenthesis are the lager group reported for that yeast (see Supplementary Table S1 for references information). SACE: S. cerevisiae; SAEU: S. eubayanus. * Study yeast.
pipeline, we aligned using BOWTIE2 (Langmead and Salzberg 2012) the BESs collection with several lager yeast assemblies of different origin, sequencing, and assembly technologies (Supplementary Table S1). These were the S. pastorianus 790 assembly (De León-Medina et al. 2016), 10 S. pastorianus assemblies (Nakao et al. 2009; Dostál et al. 2013; Walther et al. 2014; van den Broek et al. 2015; Okuno et al. 2016) (Supplementary Table S1), and the Saccharomyces sp. strain M14 assembly (Liu et al. 2018). The percentage of total aligned BESs of group-1 yeast assemblies ranged from 60.74% to 80.74% (Table 2). The range for group-2 yeast assemblies was 88.06–97.0% (Table 2). The difference in the total percent of aligned BESs among the lager groups could be due to a higher nucleotide identity of the group-2 yeast sequences with the same group yeast BESs data set. Alternatively, it could be due to the larger genome size of group-2 yeasts and, as consequence, absent sequences in group-1 yeasts. The percentage of total aligned BESs did not correlate with the N50 of the assemblies (R = 0.12, P < 0.05), which suggest that the BESs can be aligned to any S. pastorianus assembly, independently of the assemblies’ contiguity, and that the alignment results may not be biased by this assembly’s parameter. Oppositely, the percentage of paired-end BESs varied largely among assemblies but correlated with their N50 (R = 0.96, P < 0.05), which indicates that the number of this type of alignments depends on the assembly’s completeness. The percentage of the rest of the alignment types, such as unpaired-end alignments, may also be related to the N50 of the assemblies; but it could also be due to genomic differences between the strains. Therefore, the S. pastorianus 790 BESs alignments with other lager yeast assemblies only exemplify the type of results that could be obtained with BESs data, and information from strain specific BAC libraries should be used for correct interpretation. Finally, simplistic graphics of the alignments results can be obtained (Figure 2), which are intended to provide the potential users a rapid overview of the pipeline results. Similarly, a specially formatted file can be used to aid the visual inspection of the alignments results (Figure 3). Altogether, the pipeline works properly with any given S. pastorianus assembly, regardless of their sequencing and assembly technology, and BESs can be aligned with the assemblies independently of their contiguity.

**BESs alignments validate and improve the S. pastorianus 790 assembly**

To assess the BAC library information potential, we analyzed in detail the BESs alignments with the available S. pastorianus 790 assembly (De León-Medina et al. 2016). We observed that 93.31% of the BESs could be located in the 790 strain assembly, whereas 6.69% did not align (Figure 2A and Supplementary Table S4). Of the aligning BESs, 16.73% were paired, 66.83% were unpaired-end, and 9.74% were singletons (Table 3 and Figure 2B). We did not find BESs alignments in positive, negative, or opposite orientations. Only paired-end alignments are considered in agreement with biological data and, therefore, validated the covered regions (Figure 3B). Unpaired-end BESs alignments indicate potential assembly improvements, such as contig joins (Figure 3C); however, these contig joins can only be made if the hypothetical insert size is close to the one observed experimentally. Otherwise, unpaired-end BESs alignments may indicate discrepancies in the experimental data compared to the assembly, such as assembly errors, or something else. Therefore, we categorized the unpaired-end alignments according to their hypothetical insert size in greater or less than 100 kbp (the estimated rounded insert size
of the BAC library). We observed that 1.17% unpaired-end BESs had an insert size >100 kbp, and as mentioned, they point differences between the BESs and the assembly (or, to a lesser extent, presence of BESs from clones with unrelated ligated inserts). Whereas the remaining majority of the unpaired-end BESs (65.66%) had an insert size of <100 kbp, suggesting contig joins (Figure 2C). Accordingly, we found 334 regions with consistent unpaired-end BESs alignments (Figure 2D and Supplementary File S2). To proof that unpaired-end alignments with short insert size (<100 kbp) could be used to improve the assembly, we designed primers flanking three possible joins (Supplementary Table S3) and amplified the region using the DNA obtained from spanning clones as template. All three regions were successfully amplified, indicating that the information contained in the BACs agrees with the bioinformatic inferences (Figure 4). Overall, BESs can be used to validate and improve the available assembly, and BACs are a reliable source of information.

**BESs substantially improve the *S. pastorianus* 790 assembly by scaffolding**

Our previous analysis suggested that many contigs could be joined with the BESs. To test whether the long-insert information of the BAC library sequences could be used to improve the *S. pastorianus* 790 available assembly (De León-Medina et al. 2016), we used the BESs alignments with BOWTIE2 to scaffold the contigs using SSPACE (Boetzer et al. 2011). The number of sequences went from 1098 contigs in the original assembly to 1001 scaffolds in the scaffolded version. The former seemed like a modest improvement; however, more than a half of the new assembly (523 scaffolds) consisted of scaffolds smaller than 1000 bp that were already present in the initial assembly. Nonetheless, the N50 statistic improved from 85,203 bp (with 75 contigs) in the first assembly to 271,957 bp (with 28 contigs) in the last one. The later N50 is larger than the smallest chromosome of *S. cerevisiae* S288c (Engel et al. 2014), suggesting that the largest scaffolds in the current assembly are chromosome-sized. We realigned the BESs to the scaffolded sequences and we observed that the percentage of paired-end BESs alignments increased from the original 16.73% to 45.05%; whereas the percentage of unpaired-end BESs alignments decreased from 66.83% to 38.41% (Table 3), indicating a better agreement of the biological data with the new scaffolded assembly. Notwithstanding, when scanning for consistent unpaired-end BESs alignments, 147 regions still harbored potential contig joins. The former could be due to unresolved unions originated from repeated regions of assorted kinds, as well as homeologous translocations. Nevertheless, the use of BESs information in combination with scaffolding produces a substantially improved assembly with respect to the initial one.
BESs indicate the presence of homeologous translocations but not new nonhomologous translocations in the S. pastorianus genome

Previously, De León-Medina et al. (2016) observed nonhomologous translocations in the S. pastorianus 790 assembly, whereas homeologous translocations were not searched. To systematically detect nonhomologous and homeologous translocations present in the BESs collection, we assigned the BESs to a parental sub-genome by aligning them with the combined parental-like genomes of S. cerevisiae S288c (Engel et al. 2014) and S. eubayanus FM1318 (Baker et al. 2015). BESs that align in different sequences (unpaired-end) within the same parental genome are thought to harbor nonhomologous translocations, but if such BESs align in different parental genomes, then they are thought to harbor nonhomologous translocations but not new nonhomologous translocations in the S. pastorianus 790 genome.

Table 3 BESs alignments results with the original De León-Medina et al. (2016) assembly and improved assemblies’ versions, categorized by alignment type and expressed as percentages (%)

| Alignment type | Original | Scaffolded | Joined | Splitted |
|----------------|----------|-----------|--------|----------|
| Paired-end     | 16.73    | 45.05     | 65.53  | 62.89    |
| Opposite       | 0        | 0         | 0      | 0        |
| Positive       | 0        | 0         | 0.25   | 0.25     |
| Negative       | 0        | 0         | 0      | 0        |
| Unpaired-end   | 66.83    | 38.41     | 3.79   | 6.39     |
| >100 kbp       | 1.17     | 4.42      | 2.52   | 1.12     |
| <100 kbp       | 65.66    | 33.99     | 1.27   | 5.27     |
| Singletons     | 9.74     | 9.79      | 14.54  | 14.56    |
| BES total      | 93.31    | 93.25     | 84.11  | 84.09    |

BESs indicate the presence of homeologous translocations but not new nonhomologous translocations in the S. pastorianus 790 genome.

Figure 3 BESs alignments visualization with the S. pastorianus 790 De León-Medina et al. (2016) assembly. (A) Contig 1. (B) BESs alignments. (C) Unpaired-end BESs alignments scanning results. Most of the BESs alignments in contig 1 were paired-end (represented with their simulated insert as thick lines with “Paired” prefix). This type of alignments validates the assembled region. We also observed unpaired-end BESs aligned positively (right arrows with “Pos_Unp” prefix), unpaired-end BESs aligned negatively (left arrows with “Nega_Unp” prefix), and single (left arrows with “Single” prefix). The unpaired-end BESs scanning script detected two regions with consistent unpaired-end BESs representing possible contig joins to contig00062 and contig00176 (thin lines with names nega_bes_contig00062 and posi_bes_contig00176).

Table 3 BESs alignments results with the original De León-Medina et al. (2016) assembly and improved assemblies’ versions, categorized by alignment type and expressed as percentages (%)

Figure 4 Gaps and homeologous translocations amplification. Vectors were purified from clones spanning the potential contig join or homeologous translocation. Primers flanking the join were designed and used for PCR. Three contig joins (lanes 1–3) were successfully amplified. Likewise, homeologous translocations ScXI-SeXI (lane 3) and SeXIII-ScXIII (lane 6) were detected. ScXII (lane 4) and SeXII (lane 5) alleles, and a ScXIII (lane 7) allele, were also identified by PCR. Primers spanning Gap2 (lane 2) were also used to confirm the union between scaffold 17 and scaffold 22 of an S. pastorianus 790 assembly improved by scaffolding (De León-Medina et al. 2016). Primers used to amplify the union of Gap3 were also flanking the chimeric allele ScXI-SeXI (lane 3). Clone names used are followed by the underscore sign of each lane name.
homeologous translocations. The results showed that most of the BESs alignments (79.72%) in the combined parental genomes were paired-end (Supplementary Figure S5b) and all the parental chromosomes were covered by this type of alignments (except for chromosomes III and VII of the S. eubayanus genome that were partially covered), which suggest that S. pastorianus 790 possesses at least the same chromosomes of both parental yeasts. The unpaired-end BESs alignments were 5.19% (Supplementary Figure S5b) and, as established, indicate potential translocations in the S. pastorianus 790 genome. Therefore, this type of alignments were scanned for regions with consistent BESs. Of the found regions, 14 occurred between chromosomes of different parental type, which translated to seven potential recombinations between homeologous chromosomes III, VII, X, XI, XIII, and XVI in two regions (Supplementary File S3). Four additional regions, which corresponded to two potential homologous translocations, one in a second position of chromosomes X, and another in a third position of chromosomes XVI, were identified with a lower threshold (File S4) and included to the list because we considered BESs a reliable source of information, and because they were near to genes YJR009C/TDH2 and YPR191W/QCR2, other previously reported recombination sites in group-2 lager yeasts (Nakao et al. 2009; van den Broek et al. 2015; Monerawela and Bond 2017). In total, nine potential homologous translocations were identified with the S. pastorianus 790 BESs and the parental genomes as reference (Supplementary Figure S5d). To validate these observations, two possible homologous translocations in S. pastorianus 790, between chromosomes ScXI and SeXI, and chromosomes ScXIII and ScXIII, were confirmed by PCR amplification using the BACs DNA as template (Figure 4). Opposed to what De León-Medina et al. (2016) described, we did not find nonhomologous translocations in the unpaired-end alignments (File S3). However, S. eubayanus possesses two nonhomologous translocations between chromosomes II and IV, and between chromosomes VIII and XV, which were already present in this parental yeast prior to the hybridization event (Baker et al. 2015). These translocations are present in the Baker et al. (2015) assembly and did not produce unpaired-end BESs alignments, the kind of alignments we used for searching translocations. The above means that the translocations of chromosomes II and IV and VIII and XV are also present in S. pastorianus 790. To confirm that other nonhomologous translocations detected by De León-Medina et al. (2016) were inaccurate, we aligned the BESs to a potential nonhomologous translocation present in scaffold 17 of an S. pastorianus 790 assembly improved by scaffolding with short-insert sequencing libraries (De León-Medina et al. 2016) (Supplementary File S1). We observed that the region (near loci YBL074C and YER092W) was not covered by paired-end BESs alignments, which suggest that this region is not sustained by the BESs experimental evidence (Supplementary Figure S6). The region was also flanked by unpaired-end alignments pointing out a union between scaffold 17 and scaffold 22. Using BACs DNA as template, we confirmed the union of scaffold 17 and scaffold 22 by PCR (Figure 4). Furthermore, the region had a gap (Supplementary Figure S6 and Supplementary File S1) possibly generated during the De León-Medina et al. (2016) scaffolding, a procedure that has been reported introduces errors (Martin et al. 2016). The former findings indicated that the nonhomologous translocation in the De León-Medina et al. (2016) scaffold 17 is a misassembly. In summary, the S. pastorianus 790 genome structure, with its chromosomes homeologous to both parental types, is represented in the BAC library; moreover, the BESs information allows the detection of homeologous translocations between the chromosomes and also misassemblies.

**Saccharomyces pastorianus 790 genome structure is similar to that of other group-2 yeasts**

With the prior information, specifically the BES-supported scaffolded assembly version, and the parental chromosomes and homeologous translocations detected with the BESs, we proposed a hypothesis of the genome structure of S. pastorianus 790 based on the following observations. We used the parental-like genomes of S. cerevisiae S288c (Engel et al. 2014) and S. eubayanus FM1318 (Baker et al. 2015) to order and join the scaffolds into chromosomes using NUCMER (Kurtz et al. 2004). Given that ordering the scaffolds using another species reference may bias the result, we considered the output as a potential genome structure and not as a new assembly. We realigned the BESs with this genome structure, detected the site of potential homeologous translocations using our unpaired-end BESs scanning script (Supplementary Figure S7), splitted the sequences, and generated alternative chromosomes. We annotated the new genome structure and assigned the sequences to a parental type using the observed gene information. With the former, we could reconstruct 35 chromosome-sized pseudo-molecules. Of these, 12 were formed by sequences mainly containing genes of one parental type and that were present in both allelic versions (Sc and Se). These were chromosomes ScI, ScV, ScVII, ScVI, ScIX, ScIX, ScXII, ScXII, ScXIV, and ScXV (Figure 5A). Eight chromosomes almost exclusively contained genes assigned to one parental type and were present in only one parental allelic version. These were chromosomes ScII, ScIV, ScVIII, and ScXV from the S. cerevisiae sub-genome and ScII-ScV, ScIV-ScII, ScVII-ScXV, and ScXV-ScVII from the S. eubayanus sub-genome (Figure 5B). Fifteen sequences formed chromosome groups present in one or both parental versions plus one or two chimeric versions (i.e., contained long stretches of genes assigned to one or the other parental type, and hence harbored homeologous translocations). These were the chromosome groups ScIII and ScIII-ScIII; ScVII and ScVII-ScVII; ScX, ScX, ScX-ScX, and ScX-ScV; ScXI, ScXI, and ScXI-ScXI; ScXIII and ScXIII-ScXIII; and mosaic ScXVII and mosaic ScXVII (Figure 5C and Supplementary Table S5). Monerawela and Bond (2017) reported for S. pastorianus 790 a second translocation in the YGR285C-ZUO1 gene of chromosomes VII. We could locate one ZUO1 gene in the sub-telomeric region of each chromosome ScVII and ScVII-SeVII. When we inspected the BESs alignments near the ZUO1 gene of chromosome ScVII, we observed a small region (approximately of 10,000 bp) with two unpaired-end BESs alignments pointing to chromosome ScVII-SeVII (Supplementary Figure S8); therefore, we also considered this translocation in the genome structure. The S. pastorianus 790 proposed structure is highly similar to that of other lager yeasts such as S. pastorianus WS 34/70 (Supplementary Table S5) (Nakao et al. 2009; Monerawela and Bond 2017), which suggest that potential differences between S. pastorianus 790 and other yeast of the same type may be due to other genome features instead of the genome structure exclusively. Overall, it was possible to propose a genome structure of S. pastorianus 790 exploiting the information contained in the BESs.

**High copy chromosome number and ploidy levels in S. pastorianus 790**

The fact that the genome structure of our study yeast was similar to that of other group-2 lager strains prompted us to question whether there were other differentiating features in the S. pastorianus 790 genome. It is thought that group-1 lager yeast strains
are generally allotriploids, whereas, group-2 yeasts are thought to be allotetraploids (Walther et al. 2014; van den Broek et al. 2015). We measured the DNA content of S. pastorianus 790 and other yeasts with known ploidy levels. We observed that the DNA content of S. pastorianus 790 was of 60.1 Mbp, which is approximately five times larger than the S. cerevisiae S288c genome (Engel et al. 2014) (Figure 6), and suggesting that S. pastorianus 790 is an allopentaploid. To further investigate this, we took one short-reads sequencing library from a previous whole genome sequencing of S. pastorianus 790 (De León-Medina et al. 2016), and we obtained the approximate chromosomes copy number using the parental chromosomes as reference. We observed that the overall number of chromosomes ranged from one to three (Supplementary Table S6; Supplementary Figures S9 and S10), and for the specific case of chromosomes with homeologous pairs the chromosomes sum ranged from four to five (Supplementary Table S6). We also observed copy number changes in all the regions flanking the homeologous translocations detected using the BESs. These copy number changes suggest recombination sites and validate our previous inferences. The only exceptions from the former observation were chromosomes X, which exhibited no copy number changes across the sequence and an equal chromosome copy number. Finally, when taking chromosome sizes and their estimated copy number, we obtained a total size of 55.79 Mbp (Supplementary Table S6), which is close to the genome size we determined experimentally (Figure 6). All of these results favor the hypothesis that S. pastorianus 790 is an allopentaploid.
Discussion

The use of BACs and its end-sequences validates and improves the available S. pastorianus 790 assembly. Furthermore, BACs can be used to study the genome structure of this yeast. The generated library and sequences represent an independent experimental alternative that contain several levels of information, namely paired-end, long-insert (around 100 kbp), and single allele information, that are particularly advantageous compared to other strategies to study this hybrid genome with a high ploidy level (Supplementary Table S7).

BESs, as the Hi-C technique (Marie-Nelly et al. 2014), are an independent experimental resource. This is an advantage compared to other options to produce better assemblies that include bioinformatic finishing (Gordon and Green 2013; Hunt et al. 2013; Schatz et al. 2013) or different assembly strategies such as phasing (Przyszcz and Gabaldón 2016; Fay et al. 2019), because these latter options rely on in silico analyses exclusively and use the same source of information generated for the assembly. Instead, our strategy combines newly generated data with bioinformatic analyses, which provides an additional source of information to confirm observations. Furthermore, BACs of the end-sequences are preserved and used later for experimental verification (Ammiraju et al. 2005), as we showed by PCR amplification of the suggested joins and homeologous translocations, and to facilitate experiments on heterozygous regions (Barrera-Saldaña et al. 2017). The former pose an additional advantage compared to short-reads sequencing libraries that are discarded after the process (Supplementary Table S7).

The BESs pairing information can be used to improve the S. pastorianus 790 assemblies in two ways. First, unpaired-end BESs alignments <100 kbp can be used to join contigs by scaffolding with SSPACE (Boetzer et al. 2011) and substantially improve the N50 of the assembly (with scaffolds the size of chromosomes). Second, when the unpaired-end BESs alignments hypothetical insert is larger than expected (>100 kbp), these alignments detect misassemblies. The use of BESs could be relevant because despite the great availability of lager yeast assemblies (Nakao et al. 2009; Dostálek et al. 2013; Walther et al. 2014; van den Broek et al. 2015; De León-Medina et al. 2016; Okuno et al. 2016; Liu et al. 2018), many of them have a small N50, probably due to the complexity of sequencing and assembling hybrid genomes. To improve them, specific BESs collections could be generated and compared with their assemblies, independently of the N50, given that the total of aligned BESs does not correlate with this parameter. Whereas paired-end BESs alignments could validate regions and be seen as a measurement of an assemblies’ completeness, since the number of BESs in this orientation correlates with the N50.

The long-insert BESs information can be used to span long repeated regions during scaffolding. Scaffolding (Boetzer et al. 2011) is an approach that has been used before to improve lager yeast assemblies (van den Broek et al. 2015; De León-Medina et al. 2016; Okuno et al. 2016); however, it is a bioinformatic-only approach that can introduce joining errors (Martin et al. 2016). This is indeed what we observed in the misassembly of scaffold 17 of the De León-Medina et al. (2016) assembly, which was probably generated during their scaffolding and due to the inability of the short-insert libraries (around tens or less kilobases) to span repeated regions. Supporting this, the homologous region of scaffold 17 in S. cerevisiae (near loci YBL074C and YER092W) is a repeated region containing genes derived from the whole genome duplication event (Engel et al. 2014) that may have caused the joining error. Unlike mate pairs sequencing libraries commonly used for scaffolding, BESs have an insert length that can easily surpass these repeated regions (Supplementary Table S7).

Most importantly, the single allele information contained in the BESs was key to detect chimeric chromosomes in the S. pastorianus 790 genome. Many assembly algorithms retrieve one consensus sequence and do not allow to easily retrieve alternative alleles (Supplementary Table S7). Detection of chimeric chromosomes and misassemblies had to be performed in a blind automatic process (Hunt et al. 2013; Przyszcz and Gabaldón 2016) or manual in a time-consuming way not specially designed for hybrids and chimeric chromosomes (i.e., without a clear mechanism to deal with regions with paired-end and unpaired-end reads simultaneously aligned) (Gordon and Green 2013; Schatz et al. 2013). The pipeline presented in this work combines both, automatic and manual approaches, by automating the searches for our specific type of data, suggesting all possible joins or alleles, and facilitating the visual inspection of regions with misassemblies and homeologous translocations for the user to decide. Furthermore, the resulting files (in BED and GFF format) can be used to guide analyses with other more commonly used tools such as Biostrings (Pagès et al. 2019). One possible disadvantage could be the high number of chimeric regions or joins to inspect whether our approach is applied to other highly heterozygous genomes. Notwithstanding, it was possible for us to propose a genome structure of S. pastorianus 790 exploiting the single allele information contained in the BESs, which was not possible with the assembly and scaffolding solely. Another possible disadvantage of our approach could be the reduced throughput, higher cost, and time required for Sanger sequencing compared to massive parallel sequencing (Supplementary Table S7). Nevertheless, new techniques to generate BESs using short-reads sequencing have been developed (Wei et al. 2017; Yang et al. 2020), which circumvent the downsides of Sanger sequencing and make the application of BESs and approaches such as the one described in this work more feasible.

The lager genome structure has been widely studied with special focus on chimeric chromosomes (Nakao et al. 2009; Walther et al. 2014; van den Broek et al. 2015; Monerawela and Bond 2017). All the chimeric chromosomes we detected in our studied yeast have been already reported in other lager strains (Nakao et al. 2009; van den Broek et al. 2015; Monerawela and Bond 2017). For instance, the S. pastorianus 790 genome structure is very similar to that of S. pastorianus WS 34/70 (Supplementary Table S5) (Nakao et al. 2009; Monerawela and Bond 2017). Of the 36 reported chromosomes of WS 34/70 strain, we could identify 35 equivalent sequences. We did not find a chromosome with the nonhomologous translocation ScV-ScX1 or any other nonhomologous translocations in the S. cerevisiae sub-genome. Similarly, almost all the homeologous translocations we observed in S. pastorianus 790 have been detected previously in this same yeast by other researchers (Monerawela and Bond 2017), through a strategy that uses sequence similarity among known recombination sites. Nonetheless, we did not detect an introgression in chromosomes XII probably because it is small and did not have BESs aligned that would give unpaired-end alignments. Contrarily, we observed two additional translocations in chromosomes Sef1-ScIII and SeX-ScX that were not detected by the Monerawela and Bond (2017) method, probably because the recombination site was not present in the highly fragmented assembly of De León-Medina et al. (2016). Therefore, the use of BESs provides an improved detection of homeologous translocations (Supplementary Table S5). We noticed that three homeologous translocations detected with a lower threshold (YGR285C/ZUO1, YJR009C/TDH2, and
YPR191W(QCR2) were near centromeric or telomeric regions. This lower sequence representation could have occurred due to the difficulty of cloning these particular chromosomal regions. Another technique to detect homeologous translocations is to align the whole genome sequencing reads to the parental chromosomes (Hewitt et al. 2014), which we used to estimate the chromosome copy number and that in part helped us to corroborate our proposed genome structure. Nevertheless, translocations involving chromosomes with an equal copy number, such as the case of chromosomes ScX-SeX and SeX-ScX, cannot be observed by this method and became evident only when studying the unpaired-end BESs alignments, which represent and advantage of the BESs over the reads alignment strategy (Supplementary Table S7).

More intriguing was the S. pastorianus 790 ploidy level compared to other lager yeasts. The genome size of 790 strain, established as an allopolyploid, is the highest determined experimentally for this type of yeasts. This is interesting because group-2 strains formed a more compact group in the phylogenetic tree, which could be an indication of lower genetic variability of yeasts belonging to this lager group. Likewise, the genome structure of S. pastorianus 790 and other group-2 yeasts (Supplementary Table S5) appeared to be highly conserved, i.e., they shared many recombination sites. Hence, the key differences between S. pastorianus 790 and other group-2 strains may rely on the copy number of chromosomes and genes. Indeed, deviating chromosome copy number has been associated with specific phenotypes (Bolat et al. 2008; van den Broek et al. 2015). The former could be explained by a gene dose effect or by single nucleotide differences in the coding and/or regulatory sequences of the alleles. In the last case, BACs could become handy to study dissimilarities between the alternative copies, since they originated from individual DNA molecules. The specific phenotype resulting from the extraordinary S. pastorianus 790 ploidy level is yet to be determined. Nonetheless, these conjectures may give direction to future experiments. For example, genome diversity of lager yeast strains is thought to be limited due to an evolutionary bottleneck that occurred during the industrialization of lager beer production (Gallone et al. 2019). However, extra-variation and innovation in the lager beer industry could be achieved by screening for different ploidy levels yeasts in available collections or by artificially increasing the number of chromosome copies by genetic modification, physical or chemical treatments (Gorter de Vries et al. 2017, 2020).

Beyond the proposed genome structure and peculiarities of S. pastorianus 790, we would like to highlight the importance of the use of BESs, with their paired-end, long-insert, and single allele information, to improve short-reads assemblies and detect chromosome structural variants in a hybrid genome. Altogether, our results show the value of combining bioinformatic analyses with experimental information. Our strategy represents an alternative that can be generalized to study other hybrid genomes, especially if obtaining the most complete assembly of the study organism is in mind. The elucidated genome structure and ploidy levels will allow us to design future experiments, to understand their phenotypical implications and possible industrial applications.

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