A combined meta-barcoding and shotgun metagenomic analysis of spontaneous wine fermentation

Peter R. Sternes\(^1,3\), Danna Lee\(^1,4\), Dariusz R. Kutyna\(^1\) and Anthony R. Borneman\(^1,2, *\)

\(^1\)The Australian Wine Research Institute, PO Box 197, Glen Osmond, South Australia, 5064, \(^2\)Department of Genetics and Evolution, University of Adelaide, South Australia, 5000 Present address, \(^3\)Institute of Health and Biomedical Innovation, Queensland University of Technology, Woolloongabba, Queensland, Australia and \(^4\)Department of Cell and Molecular Biology, Uppsala University, Uppsala, Sweden

\(^*\)Correspondence address. Anthony R. Borneman, The Australian Wine Research Institute, PO Box 197, Glen Osmond, South Australia, 5064. Tel: +61 8 8313 6600; Fax: +61 8 8313 6601; E-mail: anthony.borneman@awri.com.au

Abstract

Wine is a complex beverage, comprising hundreds of metabolites produced through the action of yeasts and bacteria in fermenting grape must. Commercially, there is now a growing trend away from using wine yeast (Saccharomyces) starter cultures, toward the historic practice of uninoculated or “wild” fermentation, where the yeasts and bacteria associated with the grapes and/or winery perform the fermentation. It is the varied metabolic contributions of these numerous non-Saccharomyces species that are thought to impart complexity and desirable taste and aroma attributes to wild ferments in comparison to their inoculated counterparts. To map the microflora of spontaneous fermentation, metagenomic techniques were employed to characterize and monitor the progression of fungal species in 5 different wild fermentations. Both amplicon-based ribosomal DNA internal transcribed spacer (ITS) phylotyping and shotgun metagenomics were used to assess community structure across different stages of fermentation. While providing a sensitive and highly accurate means of characterizing the wine microbiome, the shotgun metagenomic data also uncovered a significant overabundance bias in the ITS phylotyping abundance estimations for the common non-Saccharomyces wine yeast genus Metschnikowia. By identifying biases such as that observed for Metschnikowia, abundance measurements from future ITS phylotyping datasets can be corrected to provide more accurate species representation. Ultimately, as more shotgun metagenomic and single-strain de novo assemblies for key wine species become available, the accuracy of both ITS-amplicon and shotgun studies will greatly increase, providing a powerful methodology for deciphering the influence of the microbial community on the wine flavor and aroma.

Keywords: wine; metagenomics; phylotyping; yeast diversity
Background

Wine is a complex beverage, comprising thousands of metabolites that are produced through the action of yeasts and bacteria in fermenting grape must. When grapes are crushed and allowed to ferment naturally, a complex microbial succession of yeasts and bacteria is generally observed. In the very early stages of fermentation, aerobic and apiculate yeasts and yeast-like fungi from genera such as Aureobasidium, Rhodotorula, Pichia, Candida, Hanseniaspora and Metschnikowia, which reside on the surface of intact grape berries or winery equipment, represent the majority of the microbiota [1]. However, most of these species, especially the aerobic yeasts, succumb early in the succession of the fermentation in response to falling oxygen levels and increasing ethanol. Mildly fermentative yeasts, such as Hanseniaspora uvarum, Candida stellata, Metschnikowia pulcherrima, Torulaspora delbrueckii, and Lachancea thermotolerans, can proliferate and survive well into the fermentation, but fall in numbers as ethanol levels increase, although it has been reported that C. stellata can survive up to 12% ethanol and complete fermentation [2–5].

Despite the vastly higher numbers of non-Saccharomyces yeasts early in the fermentation process, the major wine yeast, Saccharomyces cerevisiae, is responsible for the bulk of the ethanolic fermentation. However, S. cerevisiae is not readily isolated from intact grape berries and is therefore generally found in very low numbers at the start of fermentation [6, 7]. Regardless, due to its higher fermentative ability, growth rate, and tolerance to ethanol, S. cerevisiae supplants the various non-Saccharomyces yeasts, becoming the dominant species from mid-fermentation such that an almost monoculture of this 1 species is established by the end of fermentation.

While traditional microbiological techniques have provided important insights into the microbial succession that occurs in spontaneous fermentations, both the breadth of fermentations investigated and the depth at which individual species contributions could be resolved have been limited. Recent advances in culture-independent methods for species analysis, such as amplicon-based phylotyping (also known as meta-barcoding) and metagenomics provide a high-throughput means to analyze large numbers of microbiological samples at great depth [8]. Accordingly, these techniques are now being adapted for the study of wine fermentation, and several amplicon-based studies have been conducted to investigate vineyard and wine microbiomes [9–14]. However, despite these previous amplicon studies, there are still concerns regarding biases that may be inherent in the process due to uneven polymerase chain reaction (PCR) amplification or unequal copy number of the ribosomal repeat [9, 15]. To address some of these limitations, metagenomic techniques are being used to determine species abundance from shotgun sequencing of mixed samples. These techniques generally rely on read mapping, either to collections of curated marker genes or whole genomes, making them reliant on reference sequences that are available [16]. As yet, shotgun metagenomics has not been applied to the study of wine fermentation or to assess the accuracy of amplicon-based abundance estimates of wine fermentation.

Fungal-specific internal transcribed spacer (ITS) phylotyping was performed over 4 key fermentation stages in 5 independent commercial Chardonnay juice fermentations in triplicate. Full shotgun metagenomic sequencing was also performed for 20 of these samples. Comparison of the ITS phylotyping and shotgun data uncovered a major amplicon bias that existed for the genus Metschnikowia, providing the means to normalize other ITS datasets in which this species is abundant.

Data Description

To map the microflora of spontaneous fermentation, both amplicon-based ITS phylotyping (meta-barcoding) and shotgun metagenomics were used to assess community structure, while comparing the biases inherent in PCR-based phylotyping relative to the unbiased shotgun metagenomic sequencing. These data provide key insights into the species participating in wild wine fermentation, while also highlighting the at least 1 significant bias in the ITS analysis of wine yeasts. All sequencing data, including ITS barcoding and shotgun metagenomic sequencing, have been deposited in Genbank under the Bioproject accession number PRJNA305659. The datasets supporting the results of this article are available in the GigaDB repository [17].

Analyses and Discussion

Analysis of microbial communities in wild ferments

To study the reproducibility and applicability of performing metagenomics analyses of laboratory-scale uninoculated fermentations, 5 Chardonnay grape musts (Y1, Y2, Y3, T1, and T2), which were each destined to undergo winery-scale uninoculated fermentation (sourced from 2 different wineries), were fermented at laboratory scale in triplicate (Table 1). Fermenting musts were tracked for sugar consumption via refractometry, with samples taken for analysis at 4 key time points: D0, just after crush (11–12 Baumé [Bé]); D1, the onset of fermentation (10–11 Bé, ~90% residual sugar); D2, mid-ferment (5–6 Bé, ~50% residual sugar); and D3, nearing the end of ferment (~3 Bé, ~25% residual sugar). All ferments proceeded to dryness (0 Bé, ~5 g/l residual sugar), with sample Y1 being the fastest (12 days) and sample T1 taking the longest time (27 days) to complete fermentation.

Species abundance estimation via ITS-amplicon analysis

A total of 66 samples were analyzed comprising 6 control populations (2 different mock communities in triplicate) and 60 laboratory-scale fermentations (4 stages during ferment in triplicate for 5 different musts) (Table S1). DNA was isolated from the pelleted fraction of each must sample, with a 2-step PCR performed using sequences designed to amplify the fungal ITS region [9], while adding experiment-specific inline barcodes and appropriate adaptors for sequencing on the Illumina sequencing platform (Fig. S1). Following sequencing and barcode and adaptor trimming, 8.8 million reads were assigned across the samples (Table S1), with an average of over 100 000 reads per sample.

To consistently describe and compare the number of operational taxonomic units (OTUs) across the samples, all 8.8 million reads were first analyzed as a large single batch. Dereplication [18], OTU clustering [19], and taxonomic assignment [20] of this combined dataset resulted in the production of a single OTU table that encompassed all the OTUs from across all 66 samples. Abundance measurements of each individual dereplicated OTU from each sample were then mapped to this combined data table to derive the contribution of each experiment to the collective dataset (Table S2).

Mock control populations

Given previous concerns regarding the accuracy of ITS-amplicon profiling [9], 2 different mock control populations were assembled, in triplicate, from known numbers of cells obtained from individual cultures of 7 common wine-associated yeasts,
representing 6 different species and 5 different genera (Table 2). By comparing the results of the ITS-amplicon profiling of these samples with those expected from estimated numbers of input cells, nearly all species estimates were within 2-fold of their expected value, despite cell concentrations differing across 5 orders of magnitude (Table 2). However, the results for Metschnikowia appeared to be reproducibly overestimated in both control populations (18.6- and 10.5-fold), indicating that this species may display significant amplicon bias for the ITS region relative to the other samples used in the control populations.

**Laboratory fermentations**

ITS-amplicon analysis of laboratory-scale wild fermentations showed that there was a high degree of reproducibility between each of the 3 biological triplicates ($r^2 = 0.95 ± 0.02$) (Fig. S2). All the fermentations displayed an expected microbiological succession, beginning with a diverse and variable collection of fungi that progressively resolved into a population that was dominated by the major wine yeast *S. cerevisiae* (Fig. 1A). Multidimensional scaling (principal coordinate analysis, using Bray-Curtis distance) of the fermentations showed that while T2, Y1, and Y2 could be broadly classified as being dominated by Metschnikowia and Hanseniaspora at the D0 and D1 time points, the Y3 ferment was almost devoid of these genera, with the ferment characterized by high levels of Aureobasidium and Rhodotorula, primarily at D0 (Fig. 1B). T1, the slowest ferment, displayed a highly diverse D0 population of Rhodotorula, Cladosporium, and Aureobasidium, which progressed through a Hanseniaspora-dominated phase at D1 and finally to *S. cerevisiae* at D2/D3. These dominant species, and their progression during fermentation, are broadly similar to those found in previous studies of wine fermentation [12, 14].

The use of the fungal ITS marker also allowed for species-level assignment of many OTUs, and there were several genera
Figure 1: ITS amplicon abundance of uninoculated ferments. (A) Laboratory-scale ferments analyzing 4 fermentation time points in 5 different musts in triplicate. ITS sequences are grouped by genus and are colored-coded by their normalized abundance (reads per thousand reads). (B) Dissimilarity analysis of ITS-amplicon abundance. Triplicate samples from each time point were subjected to Bray-Curtis dissimilarity analysis. The PCA weightings of the top 30 genera are overlaid on the plot, with the size of the gray circles around each node proportional to the total abundance of each genus across all samples (no shading for nodes > 5000 counts). (C) Species-level ITS assignment for the genus Hanseniaspora. The individual abundance measurements for the 8 OTUs that comprise the g/Hanseniaspora category are shown, grouped by phylogenetic distance. Results are color-coded according to normalized abundance (reads per thousand reads).

for which more than 1 species was encountered. For example, the genus Hanseniaspora was represented by a total of 8 OTUs that could be grouped into at least 5 main species (by ITS sequence similarity; H. uvarum, H. opuntiae, H. osmophila, H. vineae, and H. guilliermondii) (Fig. 1C), with 2 species, H. uvarum and H. opuntiae, having 2 distinct OTUs representing each species, but which displayed coordinated changes in abundance across both juice and time point. For these species, this argues that either there were multiple strains of each species present in the ferments (with slightly different ITS sequences) that were responding similarly or that multiple OTU sequences were being produced per species (either due to heterogeneous ITS repeats or PCR artefacts).

Interestingly, these 2 main categories of ferments that were observed (T2, Y1 and Y2, vs T1 and Y3) did not correlate with vineyard location and/or winery (Table 1). However, the overall difference in the location of the vineyards and wineries is relatively minor, with Eden Valley and the Adelaide Hills being geographically adjacent regions in South Australia. The driver of these differences in microbial starting populations and progressions therefore remains to be determined; however, undocumented factors such as vineyard management and/or microclimate may be involved [21–24].

**Shotgun metagenomics**

While ITS-amplicon sequencing provided an in-depth analysis of variation across ferments, it has been widely accepted that the combination of ITS primer sequences, multiple rounds of PCR, and variation in the ITS repeat number can produce biases in the final abundance measurements [9, 15]. In addition, unless a second primer set is employed, bacterial species are not covered by this analysis. To explore these potential biases in more detail and to potentially provide strain-level information, shotgun metagenomics, in which total DNA is extracted and directly sequenced, was employed on a total of 20 of the samples analyzed by ITS-amplicon sequencing (Table 3).

Given that reference genome sequences exist for many wine-associated microbes, a mapping abundance strategy was used to analyze the shotgun data. A representative collection of reference genomes was therefore obtained from existing genomic resources for fungal and bacterial genera that were known or
Table 3: Shotgun metagenomic read alignment statistics

| Sample                      | Total reads | Read alignment rate* (%) |
|-----------------------------|-------------|-------------------------|
| Control mix 1 replicate A   | 18 816 478  | 97.05                   |
| Control mix 1 replicate C   | 17 883 449  | 97.11                   |
| Control mix 2 replicate A   | 20 343 317  | 97.18                   |
| Control mix 1 replicate C   | 18 138 322  | 97.91                   |
| T1 D1 replicate A           | 20 027 063  | 88.15                   |
| T1 D1 replicate B           | 21 175 617  | 90.22                   |
| T2 D1 replicate A           | 18 173 778  | 90.12                   |
| T2 D1 replicate B           | 21 705 055  | 91.02                   |
| T2 D2 replicate A           | 21 282 134  | 97.12                   |
| T2 D2 replicate B           | 21 112 818  | 96.84                   |
| T1 D1 replicate A           | 20 196 267  | 96.04                   |
| Y1 D1 replicate B           | 20 404 693  | 96.39                   |
| Y2 D1 replicate A           | 18 384 104  | 93.13                   |
| Y2 D1 replicate B           | 18 960 301  | 92.61                   |
| Y2 D3 replicate A           | 20 731 661  | 96.66                   |
| Y2 D3 replicate B           | 19 327 663  | 96.75                   |
| Y3 D1 replicate A           | 19 843 426  | 94.48                   |
| Y3 D1 replicate B           | 21 559 192  | 94.29                   |
| Y3 D3 replicate A           | 19 533 468  | 96.69                   |
| Y3 D3 replicate B           | 19 258 370  | 95.77                   |

*Sequencing reads from each sample (pre-filtered to remove grapevine matches) were aligned against the wine reference consortium (Table S3).

suspected of being wine associated (Table S3). However, attempts at aligning to a preliminary reference genome set for shotgun abundance estimation (see below) resulted in up to 15% of reads being unable to be aligned.

To determine if this was due to a lack of suitable reference genomes for key species that were represented in the shotgun data, all unaligned sequences were subsequently de novo assembled, with the resulting contigs partitioned according to their likely genus. Four genera were represented by at least 500 kb of sequence, although 3 of these Rhodosporidium, Rhodotorula, and Microbotryum, represent synonymous species and were combined and ascribed to a single major assembly product, Rhodosporidium (n = 1539, 17.7 Mb). As 3 other species of Rhodosporidium and Rhodotorula are present in the reference dataset, these contigs likely represent the genome of an additional species within this genus.

A fourth group of contigs (n = 76, 644 kb) was ascribed to Aureobasidium spp., despite the presence of a reference sequence for Aureobasidium pullulans. However, in this instance, the strong correlation in abundance values obtained across the samples for these 2 different sequences point to these de novo contigs representing regions that are not conserved in the existing A. pullulans reference sequences (Fig. S3). These sequences were subsequently added to the reference set for use in the shotgun abundance estimation.

Estimating species abundance using shotgun metagenomic sequencing

The final reference genome set comprised 851 Mb of DNA that represented a total of 51 species (45 eukaryotic, 6 prokaryotic) (Table S3). After filtering each sample for reads that matched the grapevine reference genome (<1% per sample), each filtered sample was aligned to the reference genome set. This approach resulted in most reads being able to be matched to the reference consortium, although this was highly sample dependent, with between 2% and 11% of reads not able to be aligned adequately to the reference set of sequences (Table 3). These unaligned sequences likely represent species for which an adequate reference genome was not present in the consortium, while being present at too low of an abundance to produce significant contigs during the de novo assembly of the unaligned pool.

In order to determine the potential taxonomic source of these unaligned reads, the marker-gene metagenomic classifier MetaPhlAn [25] was used to classify the remaining reads from this dataset (Table S4). This indicated 40% of the remaining reads to be of bacterial origin, 46% from Ascomycetous fungi and 13% viral. The most highly represented bacterial genera included Acetobacter (25% total bacterial reads), Curtobacterium (14%), and Lactobacillus (18%), which are all commonly associated with wine or grapes [12, 26]. Of the Ascomycete spp., half the reads were predicted to be from S. cerevisiae and likely represent mitochondrial reads (the mitochondrion was excluded from the reference genome set due to its variable copy number), with another 30% predicted to derive from an unclassified member of the family Debaryomycesaceae.

For those reads that could be matched to the reference set, estimations of species abundance were made from average read coverage values from discrete 10-kb windows across each genome (See Fig. 2A for an example dataset for Hanseniaspora spp.; a full dataset is available in Fig. S3). In addition to read depth, the average identity between each read and the reference to which it mapped was also recorded. This provided an estimate of the evolutionary distance between each genomic reference and the strains or species present in each sample. These identity values were generally above 99% for the reference genomes, but were found to be significantly lower for reference sequences including Mucor circinelloides, Pseudomonas syringae, and Hanseniaspora valbienys, suggesting that the actual species or strains present in the fermentation were significantly different than the reference used (Fig. S3).

To provide single abundance values for each reference genome in each sample, overall abundance measurements were derived from the average read depth of all 10-kb windows in each genomic sequence. An additional filter of at least 20% genome coverage was also applied to limit the effect of small numbers of windows with large coverage values, such as those derived from mis-mapping or potential small-scale horizontal transfer events from very high-abundance species against otherwise no- or low-abundance genomes (e.g., S. cerevisiae and S. paradoxus), from producing spurious abundance estimations. Using this technique, it was possible to detect the presence of 25 of the eukaryotic reference sequences and 5 prokaryotes across 5 orders of magnitude (Fig. 2B). As for the phylotyping, the major eukaryotic species that were identified include the Aureobasidium spp., Rhodotorula spp., and Hanseniaspora spp. early in the ferment progression and Saccharomyces cerevisiae late in fermentation.

Prokaryotic species, which could not be examined via the phylotyping experiments, included the malolactic wine bacterium Oenococcus oeni and various species of Lactobacillus, which are also commonly found in wine. Both O. oeni and the Lactobacillus spp. have been observed in previous microbiome projects in wine [10, 12, 14] and displayed expected increases in abundance during fermentation. In addition to the “wine-associated” bacteria, the plant pathogens Pseudomonas syringae and Tatumella ptyseos, which showed high levels of abundance early in fermentation, were also detected at high abundance; however, unlike the lactic acid bacteria, the abundance of these species declined as ferment progressed.
Figure 2: Shotgun metagenomic analysis of species. (A) Shotgun sequencing reads from each sample were mapped to the wine metagenome reference set. The total reads present in non-overlapping 10-kb windows across each genome were recorded relative to genomic location. In addition to total read number, the average identity of the reads in each window compared to the reference sequence was also calculated (id factor). For clarity and space considerations, we depict here only the abundance measures for species within the 
\textit{Hanseniaspora} genus for the 2 T2D1 and Y1D1 replicates (results for all samples are presented in Fig. S3). (B) Normalized average abundance values for each reference species in each sample. Values were normalized using total read numbers in each sample (including non-aligning reads), with final values represented per million reads in each 10-kb genomic window. (C) Bray-Curtis dissimilarity analysis of the shotgun abundance data. The weightings of each reference genome are overlaid on the plot, with the size of the gray circles around each node proportional to the total abundance of each reference genome across all of the samples (no shading for nodes >10).

Comparing the shotgun metagenomic values obtained for the 4 control experiments to those expected from the estimated numbers of input cells showed that the outcomes of the shotgun analysis were within 2-fold of each other in all but 2 cases and that they were highly correlated with the ITS results ($R^2 = 0.99$) (Table 2).

When multidimensional analysis (principal coordinate analysis [PCoA], using Bray-Curtis distances) was used to compare the shotgun samples, the presence of high amounts of 
\textit{Hanseniaspora} spp. in 3 D1 samples (T1 D1, T2 D1, and Y2 D1) largely differentiated them from the 2 remaining samples. Within the 3 D1 samples that contained high levels of 
\textit{Hanseniaspora} spp., the D1 T1 and D1 T2 samples were primarily populated by 
\textit{Hanseniaspora uvarum} while the Y2 sample contained roughly equal proportions of 
\textit{Hanseniaspora uvarum} and 
\textit{Hanseniaspora osmophila} (Fig. 2C).

Comparison of shotgun and ITS-amplicon data
In addition to comparing the control values, it was possible to extract values for comparison from the full shotgun and ITS-amplicon datasets by comparing the results from a total
of 23 comparable taxonomic identifiers that were present in both experimental types (Fig. 3). For most of these taxonomic identifiers, the normalized abundance values recorded from the shotgun and ITS-amplicon experiments were highly correlated ($R^2 = 0.93$) and differed by 2-fold or less across a dynamic range of more than 4 orders of magnitude, with accuracy diminishing at levels below 100 reads/fragments per million. For those high-abundance species that were not within a 5-fold range, the previously identified ITS-amplicon overestimation bias for Metschnikowia spp. was recapitulated, confirming that this is a bias inherent in ITS analysis for this species.

**Potential implications**

Uninoculated wine ferments represent a complex and dynamic microbial community. Metagenomics and phylotyping are now allowing for the detailed analysis of large numbers of fermentation samples, shining a light on the composition of these microbial mixtures. While the ITS phylotyping provides an accurate, high-throughput means to determine species abundance, the shotgun metagenomics uncovered at least 1 major example of amplicon bias, with the Metschnikowia spp. displaying a 10-fold over-representation. However, once biases such as these have been identified, they can be corrected in future ITS phylotyping datasets to provide a more accurate species representation. As more shotgun metagenomic and single-strain de novo assemblies for key wine species become available, the accuracy of both ITS-amplicon and shotgun studies will greatly increase. This will provide a key methodology for deciphering the influence of the microbial community on the wine flavor and aroma and how winemaking interventions may be used to shape these outcomes.

**Methods**

**Mock control populations**

To assemble control samples of known microbial composition, individual cultures of 7 wine yeasts (Table 1) were grown to stationary phase in YPD, with each culture microscopically enumerated using a hemocytometer. Specific numbers of cells from each culture were then mixed to produce 2 different control populations with proportions of individual wine yeasts that spanned several orders of magnitude (Table 1). Cell mixtures were then pelleted and DNA prepared for either phylotyping or shotgun abundance measurements as for the laboratory ferments.

**Laboratory ferments**

For each of the laboratory-scale wild ferments, 20 l of Chardonnay grape juice was obtained directly from winery fermentation tanks immediately after crushing. Each 20 l sample was then split into 3 separate 3 l glass fermenters fitted with air-locks and fermented at 20 °C with daily stirring; 50 ml samples were taken...
at D0, just after crush (11–12 Bé); D1, the onset of fermentation (10–11 Bé, ~90% residual sugar); D2, mid-ferment (5–6 Bé, ~50% residual sugar); D3, nearing the end of ferment (<3 Bé, <25% residual sugar). All ferments proceeded to dryness (0 Bé, <5 g/l residual sugar).

DNA preparation

For each sample, 50 ml of fermenting juice was centrifuged for 10 minutes at 10 000 g, washed in 20 ml PBS, re-centrifuged, and then frozen at –80°C until processed. Total DNA was extracted from washed must pellets using the PowerFood Microbial DNA Isolation Kit (Mobio).

ITS-amplicon preparation and analysis

Analysis of ITS abundance from ferment samples was performed using 2-step PCR amplification followed by next-generation amplicon sequencing (Fig. S1). First-round amplification of the ITS region was performed using the fungal-specific primers BITS (ACCTGCGGARGGATCA) and B58S3 (GAGATCTRTTGYTRAAAGTT) [9], which were modified to include both an inline barcode and Illumina adaptor sequences BITS-F1-Nxxx and BITS-R1-Nxxx (Table 4). One nanogram of DNA was used in each first-round PCR (20–30 cycles, 55°C annealing, 30-second extension, KAPA 2G Robust polymerase). Second-round amplification was performed using the Illumina adaptor sequence present in the first-round primers as an amplification target, with the remaining sequences required for dual-indexed sequencing on the Miseq platform added via overhang PCR (Fig. S1). For each sample, 2 ul of first-round PCR product was used (15 cycles, 55°C annealing, 30-second extension, KAPA 2G Robust polymerase). Following PCR, all samples were mixed into a single batch and column purified (minElute, Qiagen). ITS-amplicon pools were sequenced on the Illumina Miseq sequencing platform using 2 × 300 bp paired-end chemistry (Ramaciotti Centre for Functional Genomics, Randwick, Australia).

Following sequencing, raw sequence data were quality trimmed (Trimmomatic v. 0.22 [27], TRAILING:20 MINLEN:50) and adaptor trimmed at the 3′ end to remove ITS adaptor sequences (cutadapt 1.2.1) [28], and the individual read pairs were overlapped to form single synthetic reads (FLASH 1.2.11 [29]; –max-overlap 1000 –allow-outies). These synthetic reads were then trimmed at both the 5′ and 3′ ends to remove any remaining Illumina adaptors that were directly adjacent to the inline barcodes (cutadapt 1.2.1 [28]; -a AGATCGGAAG -g CTTCCGATCT -e 0.1 -overlap 6) and sequentially partitioned according to the specific combination of inline barcode sequences at both the 5′ and 3′ end of each synthetic read using FASTX-Toolkit (v. 0.0.13; fastx_barcode_splitter.pl –bol –mismatches 1; [30]).

To calculate the abundance of individual amplicons, the entire dataset for all of the samples was dereplicated with USEARCH ([v. 7.0.1990; -derep_fulllength, -size-out] [18] and each dereplicated OTU renamed according to the md5 checksum of the OTU sequence to provide unambiguous comparison of identical OTUs across experiments. Dereplicated OTUs were then clustered using SWARM (-z-differences 1 –fastidious [19]), with a minimum final OTU size of 10 implemented using custom scripts.

Following clustering, the likely taxonomic identity of the representative sequence of each OTU was determined using the assign_taxonomy.py module of QIIME using a modified form of the standard QIIME UNITE database in which any unclassified or unidentified sequences were removed and each ITS region was trimmed to the extent of the BITS primers used for the original ITS amplification (-t sh_taxonomy.qiime_ver6_dynamic_s_10.09.2014.txt -r refs.qiime_ver6_dynamic_s_10.09.2014.BIT.unclassified.unidentified.fasta -m uclust –uclust_similarity = 0.98 –uclust_max_accepts = 10 –uclust_min_consensus_fraction = 0.4) [20]. In addition to the edited UNITE database, OTU annotations were also performed with an augmented version of the database in which several wine-specific, manually curated reference sequences were added and 3 UNITE reference sequences that were found to have erroneous annotations were either edited or removed (Supplemental File 1).

Once the results were established for the full dataset, individual dereplicated OTUs from each sample were matched back to those of the full dataset using custom scripts to provide a directly comparable and standardized assignment of each individual experimental result within the overall dataset. Final results were assembled in QIIME tabular format using custom scripts (Table S2).

Multidimensional data analysis was performed with the R phyloseq package [31] using PCOA and Bray Curtis dissimilarity measures based upon the 30 most abundant OTUs across the samples.

Shotgun metagenomics analysis

DNA from 4 control populations and 16 fermentation samples from 2 wineries were subjected to whole-genome metagenomic sequencing. Random sequencing libraries were prepared using the Truseq nano protocol (Illumina) with a ~350 bp insert size.

Table 4: ITS amplification primers used in this study

| Primer | Sequence (Illumina adaptora | in-line barcodeb | spacerc | ITS primer sequencee |
|--------|-----------------------------|-----------------|--------|---------------------|
| BITS-F1-N701 | CCTACACGACGCTCTTCGGAGTCTTCACTCTCACTCTCGGTACGGTTGTACCTTGTTCTCGGAGATGG | TAAGGCGA | AGG | ACCTGCGGARGGATCA |
| BITS-F1-N702 | CCTACACGACGCTCTTCGGAGTCTTCACTCTCACTCTCGGTACGGTTGTACCTTGTTCTCGGAGATGG | TAAGGCGA | AGG | ACCTGCGGARGGATCA |
| BITS-F1-N703 | CCTACACGACGCTCTTCGGAGTCTTCACTCTCACTCTCGGTACGGTTGTACCTTGTTCTCGGAGATGG | TAAGGCGA | AGG | ACCTGCGGARGGATCA |
| BITS-R1-N701 | GTGACTGGAGTTCACCTGGGACGGCTCGTCACTCTCGGAGATGG | TAAGGCGA | AGG | ACCTGCGGARGGATCA |
| BITS-R1-N702 | GTGACTGGAGTTCACCTGGGACGGCTCGTCACTCTCGGAGATGG | TAAGGCGA | AGG | ACCTGCGGARGGATCA |
| BITS-R1-N703 | GTGACTGGAGTTCACCTGGGACGGCTCGTCACTCTCGGAGATGG | TAAGGCGA | AGG | ACCTGCGGARGGATCA |
| BITS-R1-N704 | GTGACTGGAGTTCACCTGGGACGGCTCGTCACTCTCGGAGATGG | TAAGGCGA | AGG | ACCTGCGGARGGATCA |

aCommon sequence allowing for clustering on the Illumina flow cell.
b8-bp variable barcode.
cVariable length spacer (0–3 bp) used to unphase conserved amplicon regions and provide higher-quality sequencing on the Illumina platform.
dSequences derived from [9].

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Sequencing libraries were then pooled and run across 3 lanes of Illumina Hiseq 2 × 100 bp chemistry (Ramaciotti Centre for Functional Genomics, Australia).

Following sequencing, raw sequence data were first filtered to limit contaminating grapevine sequences by aligning each set of sequences against the Pinot Noir grapevine genome (CAAP00000000.3) [32] using Bowtie2 v. 2.2.5 in unpaired mode [33]. In all cases, >1% of total reads were found to match the grapevine genome. All unaligned reads for which both reads in a pair failed to align to the grapevine genome were retained for further analysis.

To provide a reference sequence for read mapping, whole genome sequences were collected, where possible, from a combination of species comprising either known grape and wine microbiota (including bacteria) or other fungal species identified as being present in the fermentations analyzed in this study via ITS phylotyping (Table S3). This reference sequence was divided up into discrete windows of 10 kb using Bedtools2 (v. 2.24.0; makewindows -w 10 000) [34].

Each of the filtered shotgun datasets was then aligned to this reference set using Bowtie2 in paired-end mode, with unaligned reads saved for later analysis (-fr –maxins 1500 –no-disconcordant –no-unal –un-conc) [33]. The resultant .sam files were sorted and converted to .bam format and filtered for low-quality alignments using Samtools (v. 1.2; view –bs -q 10 — sort) [35]. For each .bam file, the total read coverage in each 10-kb reference window was calculated using Bedtools2 (v. 2.24.0; coverage -counts) [34], with the mean, median, and adjusted mean (retain mean if ≥20% of the windows in that species contained ≥1 read; otherwise mean value of 0 applied) calculated from the bed window values for each species in each sample using custom scripts. In addition to coverage values, the average identity of each mapped read was calculated for each window using custom scripts that counted the number of mismatches per read (Bowtie2 XM: tag for each read) compared to overall read length.

De novo metagenomic assembly

For the assembly of uncultivated sequences that were unrepresented in early versions of the shotgun reference collection, reads that failed to align during the shotgun metagenomic analysis were de novo assembled using SPADES [v. 3.5.0; –sc –careful] [36]. The likely taxonomic source of each contig was estimated using BLASTX (ncbi_blast-2.2.31+; -task blastx-fast -outfmt “7 target seqs”) against the non-redundant database (nr; date 14 February 2015) and extracting the taxonomic source of the best blast hit. Contigs were then partitioned according this taxonomic grouping at the genus level, with genera being manually combined where appropriate.

Comparison of shotgun and ITS phylotyping data

To compare the shotgun and ITS datasets, normalized abundance measurements for common species from both the shotgun and ITS experiments were scaled relative to a theoretical abundance of S. cerevisiae set at 100% (1 million reads per 1 million ITS fragments or 1 million shotgun reads per 1 million reads per 10-kb genomic window). This provided 2 sets of directly comparable values for downstream plotting and analysis.

Abbreviations

Bé: Baumé; ITS: ribosomal DNA internal transcribed spacer; OTUs: operational taxonomic units; PCoA: principal coordinate analysis; PCR: polymerase chain reaction.

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Availability of supporting data and materials

The datasets supporting the results of this article are available in the GigaDB repository [17]. All sequencing data, including ITS barcoding and shotgun metagenomic sequencing and the non-Saccharomyces yeast de novo genome assemblies, have been deposited in Genbank under the BioProject accession number PRJNA305659.

Figure S1. Two-step amplification of the ITS region as an Illumina-ready amplicon. First-round amplification uses BITS and B58S3 primers [9] fused to inline barcodes and adaptor sequences. Second-round amplification takes advantage of the common Illumina adapters to add Illumina indexing sequences and the P7 and P5 adapters that are required for flow cell adherence and amplification.

Figure S2. Correlation analysis of replicate ferments. Pairwise comparisons were performed between triplicate samples from the 4 fermentation time points across the 5 different juices. Raw abundance measurements were compared for all significant OTUs (>10 reads). R² values are presented for each pairwise comparison (Corr), with the density of data points indicated on the diagonal plots.

Figure S3. Shotgun metagenomic analysis of species abundance in wild fermentation via read mapping. Shotgun sequencing reads from each sample were mapped to the wine metagenome reference set. The total reads present in non-overlapping 10-kb windows across each genome were recorded relative to genomic location. In addition to total read number, the average identity of the reads in each window compared to the reference sequence was also calculated (id_factor).

S1 Table. Samples analyzed by ITS phylotyping (metabarcoding). All samples studied by ITS phylotyping are listed, including grape variety, location, and total number of reads processed.

S2 Table. ITS phylotyping abundance estimations. The results of OTU clustering are provided for all samples studied by ITS phylotyping. The results of automated taxonomy assignment (QIIME) and manual BLAST annotation are also shown, along with the final taxonomic classification used for downstream analysis (kingdom, phylum, class, order, family, genus, and species).

S3 Table. Wine reference genomes. A full list of genome sequences (and their sources) used in the wine reference consortium.

S4 Table. MetaPhlAn analysis of unaligned metagenomic reads. MetaPhlAn results for the collection of unaligned shotgun metagenomic reads.
S1 File. Customized UNITE ITS database. The custom processed UNITE ITS database used for automated QIIME taxonomy assignment.

Competing interests
The authors declare that they have no competing interests.

Author contributions
A.B. designed the experiments. P.S. and A.B performed ITS phylotyping and shotgun data analysis. D.L. performed the fermentation experiments and ITS phylotyping. D.K. produced the mock control communities. A.B and P.S prepared and drafted the manuscript.

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