RESEARCH ARTICLE

Grape microbiome as a reliable and persistent signature of field origin and environmental conditions in Cannonau wine production

Valerio Mezzasalma¹,², Anna Sandionigi¹, Ilaria Bruni¹, Antonia Bruno¹, Gianni Lovicu³, Maurizio Casiraghi¹, Massimo Labra¹*¹

1 Zooplantlab, Department of Biotechnology and Biosciences, University of Milano-Bicocca, Milan, Italy, 2 FEM2-Ambiente s.r.l., Milan, Italy, 3 Agricultural Research Agency of Sardinia (AGRIS), Sassari-Fertilia, Sassari, Italy

* massimo.labra@unimib.it

Abstract

Grape berries harbor a wide range of microbes originating from the vineyard environment, many of which are recognized for their role in the must fermentation process shaping wine quality. To better clarify the contribution of the microbiome of grape fruits during wine fermentation, we used high-throughput sequencing to identify bacterial and fungi communities associated with berries and musts of Cannonau. This is the most important cultivar-wine of Sardinia (Italy) where most vineyards are cultivated without phytochemical treatments. Results suggested that microbiomes of berries collected at four different localities share a core composition characterized by Enterobacteriales, Pseudomonadales, Bacillales, and Rhodospirillales. However, any area seems to enrich berries microbiome with peculiar microbial traits. For example, berries belonging to the biodynamic vineyards of Mamoiada were rich in Bacillales typical of manure (i.e. Lysinibacillus, Bacillus, and Sporosarcina), whereas in the Santadi locality, berries showed soil bacteria such as Pasteurellales and Bacteroidales as well as Rhodospirillales and Lactobacillales which are commonly involved in wine fermentation. In the case of fungi, the most abundant taxa were Dothioraceae, Pleosporaceae, and Saccharomycodaceae, and although the proportion of these families varied among localities, they occurred ubiquitously in all vineyards. During vinification processes performed at the same wine cellar under controlled conditions and without using any yeast starter, more than 50% of bacteria groups of berries reached musts, and each locality had its own private bacteria signature, even if Saccharomyces cerevisiae represented the most abundant fungal species. This work suggests that natural berries microbiome could be influenced by pedoclimatic and anthropologic conditions (e.g., farming management), and the fruits' microorganisms persist during the fermentation process. For these reasons, a reliable wine genotyping should include the entire holobiont (plant and all its symbionts), and bioprospecting activities on grape microbiota could lead to improved viticulture yields and wine quality.
Introduction

To date, at least 5000–8000 grape cultivars showing particular traits (grape size, shape, color, and flavor) have been selected by viticulturists [1, 2]. Despite this huge diversity, variations in environmental conditions (i.e., soil composition, water management, and climate) and fermentation processes shape the contribution of these traits and modify the quality of the resulting wines. The identification of key environmental elements involved in the regional variation of grape and wine quality characteristics is a critical feature for improving wine production in terms of consumer preference and economic appreciation [3].

Several studies showed the effects of abiotic conditions on grapevine growth and fruit development, such as UV solar radiation [4], water availability [5], and nitrogen sources [6]. At the same time, biotic factors are also involved, since *Vitis vinifera* L. naturally hosts a reservoir of microorganisms [7–9] that interact with the plant and affect both the qualitative and quantitative scale of wine production.

The occurrence and effects of regional-specific microbiota in defining wine characteristics is a more controversial issue. Experimental analyses suggest that microbes colonizing berries could significantly affect grapevine and fruit health and development [10]. However, grapevine bacteria and yeasts also contribute to shaping phenotypic characteristics, such as flavour, colour, and sugar content [11], thus influencing the winemaking process as well [12, 13].

Recently, High-Throughput DNA Sequencing techniques (HTS) have been used to characterize bacterial communities of different grapevine plant portions, such as leaves and berries [14] and to assess the provenance in terms of plant portion and farming region of some microbial groups [15, 16]. Metagenomic analyses suggested that soil serves as a primary source of microorganisms with edaphic factors influencing the native grapevine microbiome, since the microbial communities of soils from the same viticultural region are quite heterogeneous [9]. Bokulich [15] showed that *Vitis* microbial biogeography is non-randomly associated with regional, varietal, and climatic factors across multiscale viticultural zones. Moreover, in 2016, the same research team [17] suggested a strong association involving grapevine microbiota, fermentation characteristics, and wine chemical composition.

Viticulturists are aware that the ground where plants grow imparts a peculiar metabolic trait on grapes and wine; this concept is usually referred to as *terroir*. In vineyards several variables could affect grape characteristics, such as soil composition and structure, topography, climate conditions, and agricultural practices. Moreover, these conditions could also influence the plant microbiome. During vinification, other variables could act on the biotransformation of grape juice, such as the environmental conditions of the wine cellar [11, 17], as well as chemical and microbial processes involved in the fermentation processes. On the whole, wine characteristics (color, flavor, fragrance, sugar content) result from the complex interaction between abiotic and biotic elements occurring in two distinct environments: vineyards and wine cellars. To better clarify the diversity and dynamics of the microbiome belonging to these two environments and its role during wine production, we analyzed grapes and musts of cv. *Cannonau*, one of the most important black grape varieties cultivated in Sardinia (Italy).

Sardinia is the second largest Italian island located in the western Mediterranean to the south of Corsica between the Italian peninsula, Spain, and Tunisia. This island is ideally suited for viticulture [18], and it is characterized by a huge number of grapevine cultivars with different morphological and chemical characteristics [19]. Due to peculiar pedoclimatic conditions occurring in different parts the island, this model offers a great opportunity to study the relationships and changes occurring at both the environmental and grape microbiome.

In the present study, we investigated the bacterial and fungal microbiome of Cannonau berries cultivated from different localities and musts produced, under controlled conditions, at
the same wine cellar. This presents the opportunity to track the microbial community from grapes to wine cellar. Specifically, the goals of our work were to: i) evaluate the microbial diversity at the vineyard level in response to different environmental conditions and farming management (e.g., biodynamic) of Sardinian localities ii) study the dynamics of microbial diversity from the vineyard to the wine cellar to estimate the impact of field bacteria on wine must.

**Materials and methods**

**Samples collection and wine production**

Given its pedoclimatic conditions and geographical isolation, Sardinian viticulture does not demand phytochemical treatments. For this reason, the island represents a suitable area to investigate the effect of natural field characteristics on the microbiome of grapes and therefore on the resulting must. To perform our experiments, we selected four Sardinian localities: Alghero (ALG), Mamoia (MAM), Mores (MOR), and Santadi (SAN) (Fig 1).

Climatic and pedological conditions and the managed condition of vineyards for each locality are provided in Table 1.
During the harvest of 2015, mature grapes (20 degrees Brix) of the most diffused cultivar ‘Cannonau’ were collected as bunches bulks (100 berries) at each locality. One degree Brix is 1 gram of sucrose in 100 grams of solution and the scale is used as a proxy for grape maturation and fermentation progress. Sampling was performed in collaboration with specialized technicians of Agricultural Research Agency of Sardinia (AGRIS) at vineyards of the four localities. Although no specific authorization was required for sampling activities, vineyards owners gave permission to conduct the study on these sites. Field studies did not involve endangered or protected species. The collected Grape samples (G) were immediately frozen, shipped on ice, and stored at −80˚C. These samples were used to characterize the berries microbiome (i.e. bacteria and yeast communities) of each locality.

To evaluate the effect of the environmental microbial community on must, we performed a wine-making process in controlled conditions starting from mature grapes collected from each locality. This was carried out at the experimental winery of AGRIS in a scale of 100 L per locality, without wine yeast starter and sulfur dioxide treatment. These conditions provide a natural fermentation process without any forced microbial selection [20]. This strategy is an emerging trend of winemaking to enhance natural interactions between microorganisms occurring during vinification [21]. To assess the characteristics and changes of must microbiome, we identified two phases based on the analysis of fermentation curves (which are usually conducted during winemaking process), namely initial must (iM) and end must (eM). The former has been collected about six hours after pressing, when the must shows the highest level of glucose (at least 20 degrees Brix) and the lowest level of ethanol (100 ml of iM for each locality). End must has been collected at 7 days after the winemaking process has started and when glucose level is lower than 2 degrees Brix and ethanol reach 12% v/v (100 ml of iM for each locality). In the case of iM samples, we were interested in evaluating the effect of the wine cellar on the original grape microbiome, whereas in the case of eM samples, we tested the dynamics of microbiome composition during the fermentation process. Each sample was stored at −80˚C.

DNA extraction

Microbial biomass recovery from G samples was obtained starting from twenty berries randomly selected from each vineyard. These were thawed and placed in 500 mL sterile Erlenmeyer flasks. Berries were washed with 100 mL of isotonic solution (0.9% w/v NaCl) for 3 h with agitation at 150 rpm. The obtained cell suspensions were separated from the berries by centrifugation at 6,000 × g for 15 min. Pellets were stored at -20˚C until DNA isolation.

In the case of must, microbial biomass was obtained from 10 mL of iM and eM samples. These were thawed and centrifuged at 6,000 × g for 15 min, washed three times in ice-cold isotonic solution. Pellets were stored at -20˚C until DNA isolation. G, iM and eM samples were processed in duplicate.
Total genomic DNA were obtained from pellets using PowerSoil™ DNA Isolation Kit (MOLY BIO Laboratories, Carlsbad, CA, USA) following the manufacturer’s instructions with modifications specific for wet soil samples.

Before libraries preparation, the obtained genomic DNA extracts were purified using Zymo Research DNA Clean and Concentrator-10 (Zymo Research, Irvine, CA, USA) to remove PCR inhibitors.

**Library preparation and sequencing**

For each DNA sample, two independent DNA libraries, for bacteria and fungi, were prepared following Illumina guidelines (16S Metagenomic Sequencing Library Preparation, Part #15044223 Rev. B) with modifications. Bacterial V3 and V4 16S rRNA genes were amplified using primers S-D-Bact-0341-b-S-17 and S-D-Bact-0785-a-A-21 [22] with the addition of the Illumina overhang adapter sequences.

Fungal internal transcribed spacer (ITS) 1 loci were amplified with primers BITS and B58S3 [23], with the supplement of the Illumina overhang adapter sequences. Before amplification, DNA extracts were normalized by means of Quantitative real-time PCR (qPCR) Ct values with the same amplification primer pairs and the same protocols described by Bruno and colleagues [24, 25]. Finally, the obtained libraries were submitted to Polo d’Innovazione Genomica, Genetica e Biologia Società Consortile R.L. (POLO-GGB, Perugia, Italy) for Illumina paired-end library preparation, cluster generation, and 2 x 300-bp paired-end sequencing (MiSeq Reagent Kit v3) on an Illumina MiSeq instrument.

**Microbial composition and community structure analysis**

Analysis of bacterial and fungal communities were performed using scripts of the QIIME pipeline [26]. Raw Illumina reads were paired and pre-processed using USEARCH merge pairs algorithm [27]. During the Quality filter step reads were filtered out if: 1) ambiguous bases were detected, 2) lengths were outside the bounds of 250 bp and/or 3) average quality scores over a sliding window of 40 bp dropped below 25.

Bacterial reads were then processed by VSEARCH 1.1.8 software version [28], which removed noise and chimeras prior to performing de novo clustering into OTUs at 97% sequence identity and discarding those OTUs represented by less than 75 sequences. The cluster centroid for each OTU was chosen as the OTU representative sequence. The taxonomic assignment of the representative sequences was carried out using the RDP Bayesian Classifier [29] against the SILVA SSU non-redundant database (version 119 release) adopting a consensus confidence threshold of 0.8. The RDP classifier was then used for the taxonomic assignment of OTUs.

Fungal reads were cleaned concerning the noise and the chimera using the same workflow adopted for bacteria reads. Before OTUs clustering, ITSx extractor [30] was used in order to filter non-fungi contaminant reads. De novo OTUs were calculated, as in the case of bacteria, using the VSEARCH cluster algorithm at 97% sequence identity with the cluster centroid for each OTU as the OTU representative sequence. The taxonomic assignment of the representative sequences was carried out using the RDP Bayesian Classifier against UNITE fungal database [31].

For both communities, a rarefaction table was calculated for each sample to determine the suitable sequencing depth that covers the extant microbial diversity.

The intra group diversity estimation (alpha diversity) was calculated using the number of observed OTUs and the Shannon index. Community analyses (beta-diversity) were performed with qualitative (Jaccard and unweighted UniFrac for fungi and bacteria respectively) and
quantitative (Bray-Curtis and weighted UniFrac for fungi and bacteria respectively) distance metrics [32] using QIIME and phyloseq R package for statistical computing [33, 34]. Statistical significance among groups was determined by the ADONIS (permutation based ANOVA (PerMANOVA)) functions of the vegan R Package [35]. PerMANOVA Pairwise contrast was performed with R script [36].

The phylogenetic tree necessary to calculate UNIFRAC distances and based on the alignment of OTUs representative sequences was built using RAxML version 7.4.2 [37] with the GTRGAMMA model bootstrapping (1’000 replicates) best maximum likelihood tree inference. Multibar plots were generated with QIIME.

A Venn diagram was created with the online tool [38] by calculating the number of shared and unique OTUs in the different datasets.

**Results and discussion**

**Sequence analysis**

To characterize the microbial consortia associated with grapes and musts of Cannonau vineyards by HTS approach, a total of 1’600'000 and 5’000'000 quality-filtered sequences were obtained for the 16S rRNA and ITS1 marker, respectively. After the removal of low quality reads sequences failing the alignment or annotated as host or mitochondrial or chloroplast sequences, and singleton sequences, a total amount of 235’371 16S rRNA V3-V4 amplicon sequences belonging to the three fermentation steps (G, iM, eM) for 24 samples were considered for further bioinformatics analyses. These sequences had an average of 430 bp (ranging from 400 to 438 bp) with primer removal and clustered into 264 OTUs. Moreover, for the same set of samples and adopting the same procedure, a total of 216 ITS1 OTUs were supported by sequences with an average of 400 bp (ranging from 390 to 405 bp) (see for more details S1 Table).

**Bacteria and fungi OTU diversity**

OTUs diversity of bacteria and fungi was analyzed separately and described in Table 2. Considering that the two replicates did not statistically differ for each sample (R2 P v >0), we decided to combine replicates to calculate alpha diversity. Concerning bacteria, the observed

| Samples | Bacteria | Fungi |
|---------|----------|-------|
|         | Observed OTU | Shannon Index | Observed OTU | Shannon Index |
| ALG G   | 55 | 2.175450 | 110 | 2.653449 |
| ALG iM  | 59 | 2.229309 | 108 | 2.547885 |
| ALG eM  | 70 | 2.217277 | 36  | 2.281021 |
| MOR G   | 84 | 3.299958 | 117 | 2.269252 |
| MOR iM  | 94 | 1.763944 | 69  | 1.290504 |
| MOR eM  | 67 | 2.858682 | 26  | 2.464016 |
| SAN G   | 113 | 2.366787 | 133 | 1.882989 |
| SAN iM  | 103 | 2.101003 | 63  | 1.271993 |
| SAN eM  | 69 | 2.764250 | 27  | 1.971678 |
| MAM G   | 50 | 3.483344 | 94  | 1.892080 |
| MAM iM  | 46 | 2.362264 | 52  | 2.254171 |
| MAM eM  | 68 | 3.150848 | 21  | 2.391844 |

https://doi.org/10.1371/journal.pone.0184615.t002
OTUs ranged from 50–113 for G samples, 46–103 for iM, and 67–70 for eM. Data suggests that all grape samples significantly differ from each other (see data in S1 Text) with the exception of ALG-MAM (p-value = 1). This was also confirmed by Shannon Indexes (Table 2) (see data in S1 Text). The diversity among localities decreases in must samples; in the case of iM samples, only those from SAN significantly differ from ALG and MAM, but they do not differ from those of MOR (p-value = 1). Significant differences were also detected between MOR iM and MAM iM (ANOVA F = -0.5552, p = 0.00006). Finally, the OTUs diversity among localities was reduced in the case of eM samples, and no significant differences were observed among musts belonging to all localities.

In the case of fungi, all the localities share the same trend: a high number of OTUs were detected in G samples and were lower in iM and eM samples. This could suggest that several fungal OTUs occurring at fruit levels do not reach the wine cellar or do not resist the wine fermentation process as previously shown by Gilbert and colleagues [39]. Moreover, fungi species occurring in wine cellars do not largely enrich the wine microbiome in terms of OTUs diversity [23].

To better estimate the microbial dynamics of grapes from fields to the wine cellar in the four study areas, we calculated the beta diversity. Fig 2A shows the Unweighted and Weighted UniFrac PCoA plot obtained by 16S rRNA data. As previously suggested, the two sample replicates showed good overlapping (PerMANOVA pairwise test results are shown in data in S2 Text). This confirms that the sampling strategy was adequate to depict the microbiome heterogeneity of the considered vineyard. Fig 2A showed clear differences between G and musts samples. Moreover, this analysis showed that the bacterial OTUs on fruit surfaces were very different among the four localities, and this information is well represented by the unweighted non-metric multidimensional scaling NMDS (ADONIS $R^2 = 0.34$, p-value < 0.001). The iM and eM samples clustered closer. These data support the hypothesis that most of the field bacteria taxa growing on fruit surfaces are not able to persist into the wine cellar environment. This condition could be related to the fact that most of these bacteria cannot resist the change from grape aerobic to must anaerobic conditions, as well as the antimicrobial effect of secondary metabolites and ethanol occurring in must.

However, during fermentation a different bacterial community arises in must probably originating from the wine cellar. These considerations agree with previous studies [40, 41] suggesting that different microorganisms diffused in the wine cellar play peculiar roles in the specific steps of the winemaking process [11] and that their mutual interaction could affect wine characteristics.

Fig 2B shows the beta diversity of fungi OTUs. Considering the relative abundance (Bray-Curtis analysis), a pattern similar to bacteria was observed, but in this case, differences between iM and eM are more consistent. This could further suggest that wine cellar environment influences the must microbiome during the fermentation process [42]. This hypothesis is particularly supported by looking at S. cerevisiae; although this yeast was not added as a commercial starter in the 4 analyzed musts, it also occurred in iM and eM samples. The wine cellar could represent a primary source of this yeast and probably the other non-Saccharomyces yeasts, however their occurrence and development in must is related to the complex relationships among microorganisms during different fermentation phases [11, 20, 43].

To better illustrate the variation of fungi OTUs from field to wine cellar, we produced a heatmap analysis (Fig 3). This clearly showed a drop in the number of OTUs in the winery, from the initial phase of must (iM) and which became more appreciable in eM samples, as was also supported by the alpha diversity data for each locality (Table 2).
To understand the occurrence and the role of different microorganisms during the wine process and their origin, we analyzed the microbial taxonomy of the grape and must of Cannonau at different Sardinian localities.

The taxonomic assignment of sequences was performed using the RDP Classifier for both bacteria and fungi. PE sequences matching those in Silva and UNITE databases exceeding the 0.8 confidence threshold were classified and assigned to a taxonomic rank. Concerning bacteria, the 264 different OTUs were assigned to 13 predominant phyla, 44 orders, and 73 families. About 80% of OTU were identified at the genus level. In the case of fungi, a total of 216 OTUs were detected and corresponded to 4 phyla, 35 orders, and 48 families.

**Fig 2. Microbial beta diversity at the four sampling localities.** (A) Non-metric multidimensional scaling (NMDS) using unweighted and weighted UNIFRAC distance matrices as measures of beta diversity in bacterial communities. (B) NMDS using the Jaccard and Bray Curtis distance matrix coefficients to estimate the beta diversity of fungal communities. Red dots indicate grape samples (G), blue dots the initial must (IM) samples, and the green dots the end of must samples (eM).

https://doi.org/10.1371/journal.pone.0184615.g002

**Microbial taxonomy diversity**

To understand the occurrence and the role of different microorganisms during the wine process and their origin, we analyzed the microbial taxonomy of the grape and must of Cannonau at different Sardinian localities.

The taxonomic assignment of sequences was performed using the RDP Classifier for both bacteria and fungi. PE sequences matching those in Silva and UNITE databases exceeding the 0.8 confidence threshold were classified and assigned to a taxonomic rank. Concerning bacteria, the 264 different OTUs were assigned to 13 predominant phyla, 44 orders, and 73 families. About 80% of OTU were identified at the genus level. In the case of fungi, a total of 216 OTUs were detected and corresponded to 4 phyla, 35 orders, and 48 families.
Fig 4A describes the distribution of bacterial orders having a relative abundance > 0.01% determined by summing the counts derived from the two biological replicates for each sample distinguished in G, iM, and eM. Results confirm that G sample bacterial communities varied greatly across the four localities and were constituted predominantly by Enterobacteriales (19.5%), Pseudomonadales (17.5%), Bacillales (11.8%), and Rhodospirillales (8.8%). This finding agrees with data obtained by microbiome analysis performed on Grenache, one of the synonyms of Cannonau [44]. However, these four shared predominant bacteria orders should not be considered a private fingerprint of this genotype, because the same taxa were already detected on several other grapevine cultivars, such as Chardonnay, Cabernet, and Zinfandel [15]. These bacteria can be then considered a ‘common microbiome’ of vineyard soil, and they seem to not respond to pedological and environmental conditions [9, 16, 39]. The consistent presence of bacteria belonging to these orders on grape fruits could be explained by microbial migration through rain splash, winds, and insects as supported by Martins and co-workers [8], as well as by taking into account their adaptations to fruit characteristics [16]. Although the effect of these microorganisms on grape fruits and wine are unclear so far, we can conclude that a stable core microorganism of vineyards could be considered a basal biotic element able to influence different grape organs and any plant growth stages [45]. A second biotic element consists of the ‘peculiar microbial’ community that is characteristic of each vineyard and is influenced by environmental conditions and anthropogenic factors. As shown in Fig 4A, all 4
analyzed vineyards have different microbiomes at the berries level. However, samples of SAN and MAM localities showed the most peculiar microbial diversity in comparison to other localities. In SAN G samples, a consistent occurrence of Rhodospirillales (14.5%), Pasteurellales (13.1%), Bacteroidales (7.6%) was detected. Members of these taxa have been previously described in the vineyard soil microbiome. Specifically, Zarraonaindia [9] detected them in

**Fig 4. Distribution of bacterial and fungal assigned taxa.** Relative abundance of bacterial orders (A) and fungi families (B) recovered in G, iM, and eM samples at each sampling locality. Where the assignment to the Order (bacteria) or Family (fungi) rank failed, the nearest taxonomic level with supported assignment has been reported.

https://doi.org/10.1371/journal.pone.0184615.g004
five vineyards in Long Island (NY, US) characterized by granitic soil. The acidic conditions of the soil in SAN, due to the occurrence of schists-granites soil (Table 1), could support the development of similar traits to the microbiome of Long Island. Therefore, ‘peculiar’ berries microbiome could be directly influenced by soil characteristics, such as pH and soil nutritional resources [46].

Another characteristic trait of the SAN G microbiome is the modest presence of Acetobacteraceae (Rhodospirillales 14.5%) and Lactobacillaceae (1.8%), which are typically involved in the glucose fermentation of must. The occurrence of these bacteria could suggest an over-ripening of grape fruits in this vineyard, which is generally accompanied by fruit softening and partial degradation of exocarp resulting in leakage of the grape’s sugary juice. However, all sampling was performed at 20 degrees Brix, thus excluding over-ripening. Another possibility could be related to the damage of fruits. SAN samples were characterized by a relevant percentage (i.e. > 30%) of berries damaged by insects, probably belonging to Hymenoptera (AGRIS Personal communication). This event causes the fruit juice, rich in sugars content, to come out and promotes bacterial growth. In this case, external biologic factors might have influenced the berries microbiome with possible consequences on the final wine, because these microorganisms are able to actively and prematurely begin the fermentation process.

Concerning MAM samples, the collected fruits were dominated by Bacillales (32%) including members of Lysinibacillus, Bacillus (Bacillaceae), and Sporosarcina (Planococcaceae). These bacteria were found in the manure of cows, pigs, and poultry [47, 48, 49], and their occurrence in MAM could be explained by considering the biodynamic farming system applied in this vineyard. Specifically, in MAM a cow horn filled with manure was buried for maturation and subsequently was activated with water during the spring (dynamization), and the resulting product was sprayed into the field. This strategy is declared to improve soil quality, as well as enhancing plant growth and resistance to pathogens. Recent works suggested that the effect of biodynamic management could also be explained by the modification of the plant microbiome [7, 50, 51], because bacteria can act as biological disease suppression agents and could also stimulate plant growth and have an effect on plant health.

The biodynamic practice could also explain the conspicuous presence of Clostridia bacteria (4.5%) with members of Peptostreptococcaceae and Clostridiaceae. Both of these bacteria have also been detected in manure [52] and could support the beneficial effect on plant growth as well.

We conclude that in MAM the characteristic traits of the fruits microbiome are largely influenced by agricultural management and that bacteria originally related to animal manure is also able to grow on grape fruits. However on MAM G samples, we also observed members of Burkholderiales (i.e. Massilia sp.) and Rhizobiales (Rhizobium sp.) which are typically diffused in soil rich in organic matter fractions. Also, these bacteria are important players for viticulture soils, since they are able to promote plant growth [53]. On the whole, we can hypothesize that the berries microbiome characteristics reliably reflects the soil fertility of the vineyard. This is also evident in the plant of MAM vineyards, which were highly vigorous in vegetative organs, such as leaves and branches.

Concerning ALG and MOR vineyards, G samples shared the same microbial orders with a large abundance of Pseudomonadales and Enterobacteriales. This condition probably resulted from similarities in pedoclimatic characteristics: e.g. the same localities showed calcareous soil. However, some differences between these localities were detected: in MOR G samples, members of Pasteurellales were detected, whereas in ALG G samples, Xanthomonadales occurred.

Concerning the yeast diversity of G samples, the most abundant taxa were Dothioraceae (Aureobasidium, 49.86%), Pleosporaceae (Alternaria, 18.43%; Pleospora, 6.63%), and
Saccharomycodaceae (*Hanseniaspora*, 17.63%). Although the proportion of these families changed among localities, their presence was ubiquitous in all vineyards. Berries of MAM were characterized by the absence of Saccharomycodaceae, whereas the group Saccharomycetaeae was detected in SAN, but not in the others localities.

The taxonomic diversity of musts was more moderate than that of G samples with a consistent relative abundance of Rhodospirillales (from 67.2% in ALG to 50.3% in MAM), represented by the genera *Gluconobacter* and *Gluconacetobacter*, involved in the initial steps of the fermentation process. Members of Pseudomonadales, Bacillales, and Enterobacteriales occurring in G samples, dropped in musts, because they did not have the ability to grow during the wine fermentation processes [10]. Our analysis suggests moderate differences of the microbiome among samples coming from the four localities. We underline that all the collected berries were processed at the same experimental wine cellar; therefore, all musts could be influenced by the same wine cellar’s microbiome. Moreover, the winemaking process was performed under the same chemical-physical parameters. For these reasons, the only appreciable differences lay in the persistence of microorganisms deriving from the vineyard, like the Rhizobiales and Pseudomonadales (i.e., *Acinetobacter*) in MAM samples.

In eM samples, the microbiome was more variable among the samples rather than in iM samples. We detected *Gluconobacter* and *Gluconacetobacter* (Rhodospirillales) involved in must fermentation as well as *Lactobacillus* (Bacillales) involved in malolactic fermentation. In general, malolactic fermentation is most often performed shortly after the end of the alcoholic fermentation, and for this reason, it is active during eM and not in iM. In the MAM eM samples, we did not detect Lactobacillales, but there were some species described in the must, such as bacteria belonging to *Carnimonas* (Oceanospirillales) [54].

In the case of fungi, the most abundant species in must was represented by *S. cerevisiae*. Although this organism did not occur on the grape surface, starting from the first fermentation steps, it became dominant in must due to its higher fermentative ability, growth rate, and tolerance to ethanol [55]. The primary source of this yeast is crush equipment and barrel room surfaces [23], and this explains its presence in our must samples treated without any use of commercial starter. Starting from the first phases of fermentation, *S. cerevisiae* supplants the various non-*Saccharomyces* yeasts and modifies the must’s characteristics with consequences on the whole microbiome [23]. To better assess the peculiar yeasts occurring in the iM and eM samples of the four localities, the bar chart of Fig 4B was computed without *S. cerevisiae* OTUs. Data suggest that the second abundant yeast family is represented by Saccharomycodaceae with *Hanseniaspora uvarum* and other fermentative yeasts, such as members of Trichocomaceae (i.e. *Aspergillus* spp. in MOR and MAM) and Saccharomycetaceae (i.e. *Candida* spp. in SAN). Although these yeasts were less represented than *S. cerevisiae*, they are important microbial actors involved in wine fermentation, and they are able to modify wine aroma and other organoleptic characteristics through the production of a greater range of sensory-active compounds. For example, *Hanseniaspora uvarum* products 2-phenylethyl acetate, which contributes to the rose, honey, fruity and flower aromas of wines [56 57, 58]. The origin of these yeasts is partially unclear. Some of these are detected on grapes [10], while others could be resident microorganisms of wine cellar [42]. Considering that in our study all the wine juices were fermented in the same wine cellar, we could hypothesize that differences in yeasts occurring in musts could derive from the field. Although these yeasts were not detected on the berries surfaces (e.g., *Hanseniaspora uvarum, Aspergillus* spp. and *Candida*), we cannot exclude their presence as spores not detectable by our NGS sequencing analysis, but they may become appreciable in must where they are able to germinate and proliferate. Nowadays, it is critical not only to define which microbes contribute to create a high-quality wine, but also how their metabolisms can influence wine organoleptic characteristics. Integrated databases, based on
HTS and biochemical data, will permit in the very next future to analyze in depth the effects of a certain microbiome on metabolome. For example, the WineSeq platform (Biome Makers, Inc.) [59] revealed that the occurrence in G samples of some yeasts could have important effects on wine quality and human health. This is the case of the detected *Hanseniaspora* sp. yeast is related to a potential sensory profile enhancement on wine flavour.

From field to wine cellar

Many studies suggest that the grapevine’s microbiome influences the plant’s physiology, and it can then also determine some aspects of the secondary metabolites’ profile shown by fruits [11, 17]. A challenging issue to address is the impact of the berries’ indigenous microbiota on wine fermentation and the consequent effect on the sensory complexity of wines [20]. Yeasts play important roles during the alcoholic fermentation step and also have a significant impact on wine quality. However, in our experimental conditions, bacteria represented the most relevant elements of differentiation among localities (108 private OTUs out of 176 detected on G samples); therefore, we studied the persistence of the characteristic field microbiome of different localities in must samples. We underline that although bacteria are not the main driving force shaping wine characteristics and quality, they do have a significant effect on the final product. For example, lactic acid bacteria are known to convert L-malic acid to lactic acid through MLF and to impart flavor complexity, while acetic acid bacteria produce acetic acid, which is a key factor in wine spoilage. Similarly, we can expect that the bacteria occurring on Cannonau berries at different localities may play a key role in must fermentation and wine quality. However, only the investigation of bacteria communities’ dynamics during Cannonau fermentation could assess the influence of each microbial group on wine characteristics.

A Venn graph (Fig 5) shows that more than 50% of G OTUs reach the wine cellar in all localities. As suggested by Bokulich [17], vineyard-specific microbial signatures diminished during fermentation (Figs 4A, 4B and 5) as the growth of fermentative organisms reshaped the community structure, richness, and diversity of the wines. However, our data showed that more than 50% of shared OTUs between G and iM persisted in the eM phase. These bacteria could modify wine traits [11, 17, 60] not only at the field level, but also by active metabolism in must [11, 46, 53]. Fig 6 shows that those OTUs shared by field and wine cellar environments (Venn center diagram) correspond to several bacterial orders and most of these are shared by two localities at least (Fig 6). In any case, each locality has its own private OTUs. Among the investigated localities, SAN showed the highest number of unique microbial traits (see S2 Table) and representatives of Enterobacteriales, Pasteurellales, Rhodospirillales, and Lactobacillales reach the wine cellar. As previously discussed, some of these bacteria are most active during fermentation, such as *Gluconobacter* (Rhodospirillales) and *Lactobacillus* (Lactobacillales). A few OTUs were detected as unique fingerprints of ALG and MAM, but some of these were very active in must fermentation, such as the *Acetobacter* (Rhodospirillales) detected in MAM. Finally, a microbial fingerprint of MOR grape and must was characterized by several orders including the member of Caulobacteriales and Clostridiales with genera involved in plant growth stimulation [61]. Their role in wine fermentation is still unclear, but we cannot exclude that these bacteria could be able to modify some wine’s metabolites. In conclusion, we can affirm that microbiomes found at the four investigated Sardinian localities can have an impact on fruit and, must. In addition, metabolism of different microorganisms could positively enhance some wine flavor traits, but also emphasize negative organoleptic characteristics.
Conclusions

In this study, we demonstrated a distinct microbial composition of Cannonau fruits from different Sardinian localities with consequent effects also on the musts’ microbiome. To date, the role of grapevine microbes in the field has been largely ignored, with the only exception of microbial pathogens, mainly because the available technologies did not exist, and this prevented examining the community structure of the multitudes of bacterial and fungal species associated with each plant at any real depth or breadth [39]. Thanks to the HTS approach, we can now evaluate the microbial community of the grape and wine also in response to different

**Fig 5. Venn diagram of shared OTUs.** The figure shows the number of shared OTUs among sample typologies belonging to the four sampling localities.

https://doi.org/10.1371/journal.pone.0184615.g005
environmental conditions and farming practices [46]. Probably, in the very near future, this technology will be also used to deeply investigate viruses and phytoplasma that largely influence vineyard sector [62].

Emerging work clearly show that pedoclimatic conditions could affect wine characteristics not only due to the abiotic characteristics (i.e. soil, sun exposition), but also at the grape microbiome that is able to influence plant growth and development [10]. Bokulich [15] demonstrated the existence of regional microbiome fingerprints in California vineyards; Portillo [44] showed that several environmental variables, such as vineyard altitude and the geographical orientation of the plant could also affect the grape microbiome. Our study confirms that pedoclimatic characteristics could modify the fruit microbiome and underlines that agricultural practices, such as biodynamics, as well as the occurrence of opportunistic insects, such as hymenopterans, can have a consistent effect on the bacterial communities of berries and corresponding must. These results suggest that the role of the field environmental microbiome is not limited to promoting grape fruit maturation and enhancing the occurrence of some secondary metabolites strictly related to wine color and flavor, but it is also an important source of microorganisms that are able to influence wine fermentation and metabolic composition.

![Barplot showing the distribution of unique bacterial OTUs (y axis) in G samples.](https://doi.org/10.1371/journal.pone.0184615.g006)
Characteristics of the cultivars’ genotype play an important role in viticulture, so starting from the 1990’s, DNA fingerprinting approaches were used to identify synonymous cultivars and to unmask incorrect attributions. Our team [63], demonstrated a complete genetic identity between Cannonau and Spanish Grenache by using SSR markers. However, the microbiomes of these cultivars are very different [44], and appreciable differences were also observed among Grenache localities. This finding suggests that the value of the cultivar genotype is somewhat relative. A reliable genotyping should include the entire holobiont (i.e. the plant and all its symbionts [64]) of a specific Cannonau or Garnacha plant. The study of the grapevine microbiome does not represent a simple element of the product’s traceability and identity. We should consider that bioprospecting activities on grape microbiota could led to the discovery of several species with positive enological properties, as recently documented by the WineSeq® project [59]. Occurrence and abundance of these species could be easily monitored by using conventional cultivation strategies and target PCR approaches (Real-Time and Digital PCR) to be used for improving wine quality, to enhance immune capability, and reduce the use of agrochemicals. Nowadays, only an exhaustive knowledge about the vineyard, the winery and their inhabitants could permit real advancements in management activities aimed towards a better sustainable system without any loss in terms of yields and product quality.

Supporting information
S1 Text. ANOVA result on alpha diversity values for 16S and ITS1. The ANOVA and Pair-wise Post-hoc test results performed for each locality considering the three fermentation steps. (DOCX)

S2 Text. ADONIS test results. Results of ADONIS test performed to explore beta diversity patterns. (DOCX)

S1 Table. Run performance detail Table. (DOCX)

S2 Table. Table of unique OTUs distribution. Dataframe of unique OTUs distribution for localities based on Venn diagrams analysis. (XLSX)

Acknowledgments
This work was supported by the project A.K.I.N.A.S. (CUP B81H11001320002).

Author Contributions
Conceptualization: Valerio Mezzasalma, Maurizio Casiraghi, Massimo Labra.
Data curation: Anna Sandionigi.
Formal analysis: Anna Sandionigi, Ilaria Bruni.
Funding acquisition: Valerio Mezzasalma, Gianni Lovicu.
Investigation: Valerio Mezzasalma, Ilaria Bruni, Antonia Bruno.
Methodology: Valerio Mezzasalma, Antonia Bruno.
Project administration: Massimo Labra.
Resources: Gianni Lovicu.
Software: Anna Sandionigi, Antonia Bruno.

Supervision: Gianni Lovicu, Massimo Labra.

Validation: Valerio Mezzasalma, Antonia Bruno, Maurizio Casiraghi.

Visualization: Valerio Mezzasalma, Anna Sandionigi.

Writing – original draft: Valerio Mezzasalma, Anna Sandionigi, Antonia Bruno, Massimo Labra.

Writing – review & editing: Valerio Mezzasalma, Anna Sandionigi, Massimo Labra.

References

1. Fossati T, Labra M, Castiglione S, Failla O, Scienza A, Sala F. The use of AFLP and SSR molecular markers to decipher homonyms and synonyms in grapevine cultivars: the case of the varietal group known as ‘Schiave’. TAG Theoretical and Applied Genetics. 2001 Feb 1; 102(2):200–5.

2. De Mattia F, Imazio S, Grassi F, Doulati Baneh H, Scienza A, Labra M. Study of genetic relationships between wild and domesticated grapevine distributed from middle east regions to European countries. Rendiconti Lincei. 2008 Sep 1; 19(3):223–40.

3. Van Leeuwen C, Seguin G. The concept of terroir in viticulture. Journal of Wine Research. 2006 Apr 1; 17(1):1–0.

4. Fernandes de Oliveira A, Nieddu G. Accumulation and partitioning of anthocyanins in two red grape cultivars under natural and reduced UV solar radiation. Australian journal of grape and wine research. 2016 Feb 1; 22(1):96–104.

5. Martínez-Lüscher J, Morales F, Delrot S, Sánchez-Díaz M, Gomès E, Aguirreolea J, et al. Characterization of the adaptive response of grapevine (cv. Tempranillo) to UV-B radiation under water deficit conditions. Plant Science. 2015 Mar 31; 232:13–22. https://doi.org/10.1016/j.plantsci.2014.12.013 PMID: 25617319

6. Bell SJ, Henschke PA. Implications of nitrogen nutrition for grapes, fermentation and wine. Australian Journal of Grape and Wine Research. 2005 Oct 1; 11(3):242–95.

7. Compat S, Mitter B, Coll-Mull JG, Gang H, Sessitsch A. Endophytes of grapevine flowers, berries, and seeds: identification of cultivable bacteria, comparison with other plant parts, and visualization of niches of colonization. Microbial ecology. 2011 Jul 1; 62(1):188–97. https://doi.org/10.1007/s00248-011-9883-y PMID: 21625971

8. Martins G, Lauga B, Miot-Sertier C, Mercier A, Lonvaud A, Souls ML, et al. Characterization of epiphytic bacterial communities from grapes, leaves, bark and soil of grapevine plants grown, and their relations. Plos one. 2013 Aug 30; 8(6):e73013. https://doi.org/10.1371/journal.pone.0073013 PMID: 24023666

9. Zarraonaindia I, Owens SM, Weisenhorn P, West K, Hampton-Marcell J, Lax S, et al. The soil microbiome influences grapevine-associated microbiota. MBio. 2015 May 1; 6(2):e02527–14. https://doi.org/10.1128/mBio.02527-14 PMID: 25805735

10. Barata A, Malfeito-Ferreira M, Loureiro V. The microbial ecology of wine grape berries. International journal of food microbiology. 2012 Feb 15; 153(3):243–59. https://doi.org/10.1016/j.ljfoodmicro.2011.11.025 PMID: 22189021

11. Belda I, Ruiz J, Esteban-Fernández A, Navascués E, Marquina D, Santos A, et al. Microbial Contribution to Wine Aroma and its Intended Use for Wine Quality Improvement. Molecules. 2017 Jan 24; 22(2):189.

12. Capozzi V., Garofalo C., Chiriatti M. A., Greco F., & Spano G. (2015). Microbial terroir and food innovation: the case of yeast biodiversity in wine. Microbiological research, 181, 75–83. https://doi.org/10.1016/j.micres.2015.06.005 PMID: 26321127

13. Garofalo C., El Khoury M., Lucas P., Bely M., Russo P., Spano G., & Capozzi V. (2015). Autochthonous starter cultures and indigenous grape variety for regional wine production. Journal of applied microbiology, 118(6), 1395–1408. https://doi.org/10.1111/jam.12789 PMID: 25752218

14. Leveau JH, Tech JJ. Grapevine microbiomics: bacterial diversity on grape leaves and berries revealed by high-throughput sequence analysis of 16S rRNA amplicons. International Symposium on Biological Control of Postharvest Diseases: Challenges and Opportunities 905 2010 Oct 25 (pp. 31–42).

15. Bokulich NA, Thorngate JH, Richardson PM, Mills DA. Microbial biogeography of wine grapes is conditioned by cultivar, vintage, and climate. Proceedings of the National Academy of Sciences. 2014 Jan 7; 111(1):E139–48.
16. Pinto C, Gomes AC. Vitis vinifera microbiome: from basic research to technological development. BioControl. 2016 Jun 1; 61(3):243–56.

17. Bokulich NA, Collins TS, Masarweh C, Allen G, Heymann H, Ebeler SE, et al. Associations among Wine Grape Microbiome, Metabolome, and Fermentation Behavior Suggest Microbial Contribution to Regional Wine Characteristics. mBio. 2016 Jul 6; 7(3):e00631–16. https://doi.org/10.1128/mBio.00631-16 PMID: 27302757

18. Zecca GA, De Mattia F, Lovicu G, Labra M, Sala F, Grassi F. Wild grapevine: silvestris, hybrids or cultivars that escaped from vineyards? Molecular evidence in Sardinia. Plant Biology. 2010 May 1; 12(3): 558–62. https://doi.org/10.1111/j.1438-8677.2009.00226.x PMID: 20522194

19. De Mattia F, Imazio S, Grassi F, Lovicu G, Tardáguila J, Failla O, et al. Genetic characterization of Sardinia grapevine cultivars by SSR markers analysis. OENO One. 2007 Dec 31; 41(4):175–84.

20. Ciani M, Comitini F, Mannazzu I, Domizio P. Controlled mixed culture fermentation: a new perspective on the use of non-Saccharomyces yeasts in winemaking. FEMS Yeast Research. 2010 Mar 1; 10(2):123–33. https://doi.org/10.1111/j.1567-1364.2009.00579.x PMID: 19807789

21. Ciani M, Capece A, Comitini F, Canonico L, Sisto G, Romano P. Yeast interactions in inoculated wine fermentation. Frontiers in microbiology. 2016; 7.

22. Klindworth A, Pruesse E, Schweer T, Peplies J, Quast C, Horn M, et al. Evaluation of general 16S ribosomal RNA gene PCR primers for classical and next-generation sequencing-based diversity studies. Nucleic acids research. 2012 Aug 28:gks808.

23. Bokulich NA, Ohta M, Richardson PM, Mills DA. Monitoring seasonal changes in winery-resident microbiota. PLoS One. 2013 Jun 19; 8(6):e66437. https://doi.org/10.1371/journal.pone.0066437 PMID: 23840468

24. Bruno A, Sandionigi A, Galimberti A, Siani E, Labra M, Cocuzza C, et al. One step forwards for the routine use of high-throughput DNA sequencing in environmental monitoring. An efficient and standardizable method to maximize the detection of environmental bacteria. MicrobiologyOpen. 2016 Oct 1.

25. Bruno A, Sandionigi A, Rizzi E, Bernasconi M, Vicario S, Galimberti A, et al. Exploring the under-investigated “microbial dark matter” of drinking water treatment plants. Scientific Reports. 2017; 7.

26. Caporaso JG, Kuczynski J, Stombaugh J, Bittinger K, Bushman FD, Costello EK, et al. QIIME allows analysis of high-throughput community sequencing data. Nature methods. 2010 May 1; 7(5):335–6. https://doi.org/10.1038/nmeth.f.303 PMID: 20383131

27. Edgar RC. Search and clustering orders of magnitude faster than BLAST. Bioinformatics. 2010 Oct 1; 26(19):2460–1. https://doi.org/10.1093/bioinformatics/btp461 PMID: 20709691

28. Rognes T, Flouri T, Nichols B, Quince C, Mahé F. VSEARCH: a versatile open source tool for metagenomic data analysis. PeerJ. 2016 Oct 18; 4:e2584. https://doi.org/10.7717/peerj.2584 PMID: 27781170

29. Cole JR, Chai B, Farris RJ, Wang Q, Kulam SA, McGarrell DM, et al. The Ribosomal Database Project (RDP-II): sequences and tools for high-throughput rRNA analysis. Nucleic acids research. 2005 Jan 1; 33(suppl 1):D294–6.

30. Bengtsson-Palme J, Ryberg M, Hartmann M, Branco S, Wang Z, Godhe A, et al. One step forwards for the routine use of high-throughput DNA sequencing in environmental monitoring. An efficient and standardizable method to maximize the detection of environmental bacteria. MicrobiologyOpen. 2016 Oct 1.

31. Abarenkov K, Henrik Nilsson R, Larsson KH, Alexander LJ, Eberhardt U, Erland S, et al. The UNITE database for molecular identification of fungi—recent updates and future perspectives. New Phytologist. 2010 Apr 1; 186(2):281–5. https://doi.org/10.1111/j.1469-8137.2009.003160.x PMID: 20409185

32. Lozupone C, Lladser ME, Knights D, Stombaugh J, Knight R. UniFrac: an effective distance metric for microbial community comparison. The ISME journal. 2011 Feb 1; 5(2):169. https://doi.org/10.1038/ismej.2010.133 PMID: 20827291

33. Pinheiro J, Bates D, DebRoy S, Sarkar D. R Development Core Team. nlme: Linear and nonlinear mixed effects models. 2012. http://CRAN.R-project.org/package=nlme. R package version. 2012:3–1.

34. McMurdie PJ, Holmes S. phyloseq: an R package for reproducible interactive analysis and graphics of microbiome census data. PloS one. 2013 Apr 22; 8(4):e61217. https://doi.org/10.1371/journal.pone.0061217 PMID: 23630581

35. Oksanen J, Kindt R, Legendre P, O’Hara B, Stevens MH, Oksanen MJ, et al. The vegan package. Community ecology package. 2007 Oct 3; 10:631–7.

36. Arbuzu PM. How can I do PerMANOVA pairwise contrasts in R? ResearchGate. 2016 Jan 30. https://www.researchgate.net/post/How_can_I_do_PerMANOVA_pairwise_contrasts_in_R

37. Stamatakis A. RAxML version 8: a tool for phylogenetic analysis and post-analysis of large phylogenies. Bioinformatics. 2014 May 1; 30(9):1312–3. https://doi.org/10.1093/bioinformatics/btu033 PMID: 24451623
38. Genomics BE. Calculate and draw custom Venn diagrams. http://bioinformatics.psb.ugent.be/webtools/Venn/
39. Gilbert JA, van der Leie D, Zarraonaindia I. Microbial terroir for wine grapes. Proceedings of the National Academy of Sciences. 2014 Jan 7; 111(1):5–6.
40. du Toit M, Engelbrecht L, Lerm E, Krieger-Weber S. Lactobacillus: the next generation of malolactic fermentation starter cultures—an overview. Food and Bioprocess Technology. 2011 Aug 1; 4(6):876–906.
41. Lasik M. The application of malolactic fermentation process to create good-quality grape wine produced in cool-climate countries: a review. European Food Research and Technology. 2013 Dec 1; 237(6):843–50.
42. Santamaría P, Garrio P, López R, Tenorio C, Gutiérrez AR. Analysis of yeast population during spontaneous alcoholic fermentation: effect of the age of the cellar and the practice of inoculation. International journal of food microbiology. 2005 Aug 15; 103(1):49–56. https://doi.org/10.1016/j.ijfoodmicro.2004.11.024 PMID: 16084265
43. Jolly J, Augustyn OP, Pretorius IS. The role and use of non-Saccharomyces yeasts in wine production. South African Journal for Enology and Viticulture. 2006; 27(1):15.
44. Portillo MD, Franquès J, Araque I, Reguant C, Bordons A. Bacterial diversity of Grenache and Carignan grape surface from different vineyards at Priorat wine region (Catalonia, Spain). International journal of food microbiology. 2016 Feb 16; 219:56–63. https://doi.org/10.1016/j.ijfoodmicro.2015.12.002 PMID: 26736065
45. Pinto C, Pinho D, Sousa S, Pinheiro M, Egas C, Gomes AC. Unravelling the diversity of grapevine microbiome. PLoS One. 2014 Jan 16; 9(1):e85622. https://doi.org/10.1371/journal.pone.0085622 PMID: 24454903
46. Burns KN, Bokulich NA, Cantu D, Greenhut RF, Kluepfel DA, O’Geen AT, et al. Vineyard soil bacterial diversity and composition revealed by 16S rRNA genes: Differentiation by vineyard management. Soil Biology and Biochemistry. 2016 Dec 31; 103:337–48.
47. Green SJ, Michel FC, Hadar Y, Minz D. Similarity of bacterial communities in sawdust-and straw-amended cow manure composts. FEMS microbiology letters. 2004 Apr 1; 233(1):115–23. https://doi.org/10.1111/j.1567-1364.2004.00427.x PMID: 15043877
48. Lovanh N, Cook KL, Rothrock MJ, Miles DM, Sistiani K. Spatial shifts in microbial population structure within poultry litter associated with physicochemical properties. Poultry science. 2007 Sep 1; 86(9):1840–9. PMID: 17704369
49. Faissal AZ, Ouazzani N, Parrado JR, Dary M, Manyani H, Morgado BR, et al. Impact of fertilization by natural manure on the microbial quality of soil: Molecular Approach. Saudi Journal of Biological Sciences. 2017 Jan 9.
50. Bulgari D, Casati P, Brusetti L, Quaglino F, Brasca M, Daffonchio D, et al. Endophytic bacterial diversity in grapevine (Vitis vinifera L.) leaves described by 16S rRNA gene sequence analysis and length heterogeneity-PCR. The Journal of Microbiology. 2009 Aug 1; 47(4):393–401. https://doi.org/10.1007/s12275-009-0082-1 PMID: 19763412
51. West ER, Cother EJ, Steel CC, Ash GJ. The characterization and diversity of bacterial endophytes of grapevine. Canadian Journal of Microbiology. 2010 Mar 17; 56(3):209–16. https://doi.org/10.1139/w10-004 PMID: 20453907
52. Ding GC, Radl V, Schloter-Hai B, Jechalke S, Heuer H, Smalla K, et al. Dynamics of soil bacterial communities in response to repeated application of manure containing sulfadiazine. PLoS One. 2014 Mar 26; 9(3):e92958. https://doi.org/10.1371/journal.pone.0092958 PMID: 24671113
53. Burns KN, Kluepfel DA, Strauss SL, Bokulich NA, Cantu D, Steenwerth KL. Vineyard soil bacterial diversity and composition revealed by 16S rRNA genes: differentiation by geographic features. Soil Biology and Biochemistry. 2015 Dec 31; 91:232–47.
54. Pinto C, Pinho D, Cardoso R, Custódio V, Fernandes J, Sousa S, et al. Wine fermentation microbiome: a landscape from different Portuguese wine appellations. Frontiers in microbiology. 2015 Sep 1; 6:905. https://doi.org/10.3389/fmicb.2015.00905 PMID: 26388852
55. Caporaso JG, Lauber CL, Walters WA, Berg-Lyons D, Lozupone CA, Turnbaugh PJ, et al. Global patterns of 16S rRNA diversity at a depth of millions of sequences per sample. Proceedings of the National Academy of Sciences. 2011 Mar 15; 108(Supplement 1):4516–22.
56. Fleet GH. Wine yeasts for the future. FEMS Yeast Research. 2008 Nov 1; 8(7):979–95. https://doi.org/10.1111/j.1567-1364.2008.00427.x PMID: 18793201
57. Patrignani F, Montanari C, Serrazanetti DI, Braschi G, Vernocchi P, Tabanelli G, et al. Characterisation of yeast microflora, chemical and sensory properties of organic and biodynamic Sangiovese red wines. Annals of Microbiology. 2017 Jan 1; 67(1):99–109.
58. Swiegers JH, Pretorius IS. Yeast modulation of wine flavor. Advances in applied microbiology. 2005 Dec 31; 57:131–75. https://doi.org/10.1016/S0065-2164(05)57005-9 PMID: 16002012

59. Belda I, Palacios A, Fresno J, Ortega H, Acedo A. WineSeq®: A new tool for the study of the functional biodiversity of soils, and its use as a biomarker and guide for vitiviniculture practices. InBIO Web of Conferences 2017 (Vol. 9, p. 01012). EDP Sciences.

60. Styger G, Prior B, Bauer FF. Wine flavor and aroma. Journal of industrial microbiology & biotechnology. 2011 Sep 1; 38(9):1145.

61. Rana A, Saharan B, Joshi M, Prasanna R, Kumar K, Nain L. Identification of multi-trait PGPR isolates and evaluating their potential as inoculants for wheat. Annals of microbiology. 2011 Dec 1; 61(4):893–900.

62. Meng B, Martelli GP, Golino DA, Fuchs M. Grapevine Viruses: Molecular Biology, Diagnostics and Management. Springer International Publishing; 2017.

63. De Mattia F, Lovicu G, Tardaguila J, Grassi F, Imazio S, Scienza A, et al. Genetic relationships between Sardinian and Spanish viticulture: the case of ‘Cannonau’ and ‘Garnacha’. The Journal of Horticultural Science and Biotechnology. 2009 Jan 1; 84(1):65–71.

64. Vandenkoonhuyse P, Quaiser A, Duhamel M, Le Van A, Dufresne A. The importance of the microbiome of the plant holobiont. New Phytologist. 2015 Jun 1; 206(4):1196–206. https://doi.org/10.1111/nph.13312 PMID: 25655016