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Cultivable microorganisms associated with honeys of different geographical and botanical origin



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ABSTRACT

In this study, the composition of the cultivable microbial populations of 38 nectar honey and honeydew honey samples of different botanical and geographical origin were assessed. After growth in specific media, various colonies with different appearance were isolated and purified before phenotypic (morphological, physiological and biochemical traits) and genotypic [randomly amplified polymorphic DNA (RAPD), repetitive DNA elements-PCR (rep-PCR) and restriction fragment length polymorphism (RFLP]] differentiation. The identification was carried out by 16S rRNA gene sequencing for bacteria and, in addition to RFLP, by sequencing the D1/D2 region of the 26S rRNA gene for yeasts and the 5.8S-ITS rRNA region for filamentous fungi. The results showed the presence of 13 species of bacteria, 5 of yeasts and 17 of filamentous fungi; the species most frequently isolated were *Bacillus amyloliquefaciens*, *Zygosaccharomyces mellis* and *Aspergillus niger* for the three microbial groups, respectively. The highest microbial diversity was found in multifloral honeys. No correlation among the microbial species and the botanical/geographical origin was found, but some strains were highly adapted to these matrices since they were found in several samples of different origin.

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1. Introduction

According to the European Union Legislation (DLgs 179/2004) and the *Codex Alimentarius* (CODEX STAN 12-1981) honey is the natural sweet substance produced by *Apis mellifera* L bees from the nectar of plants, secretions of living parts of plants, or excretions of plant-sucking insects on the living parts of plants, which the bees collect, transform by combining with specific substances of their own, deposit, dehydrate, store and leave in the honeycomb to ripen and mature.

Honey is the most ancient sweetener used by mankind, appreciated throughout the word, embraced by religious and cultural beliefs and today considered not only a food sources, but also a homeopathic treatment alternative for wounds, burns, oral healthcare and even a potential help in cancer treatment (Lay-Flurrie, 2008; Bardy et al., 2008). It is a super saturated sugar solution characterized by a low water activity to support microbial growth (Malika et al., 2004). The natural acidity of this product, the low protein content and the high viscosity, that limit the atmospheric oxygen penetration, are particularly stressing for several microorganisms.

Honeys also possess antimicrobial properties due to several components such as glucose oxidase (Bogdanov et al., 2008), flavonoids, phenolic derivatives (Ceauşi et al., 2009) and 3-phenyllactic acid (2-hydroxy-3-phenylpropanoic acid or β -phenyllactic acid) (PLA) active against bacteria (Ohhira et al., 2004), yeasts (Schwenninger et al., 2008) and a wide range of mould species, including some mycotoxigenic species (Lavermicocca et al., 2003). However, these beneficial effects may vary depending on the product origin (Voidarou et al., 2011).

Despite the numerous inhibiting factors, some microorganisms can survive in honey, at least as latent forms and may represent a mean for their transfer to consumers. Snowdon and Cliver (1996) showed that different microbial species in honey may reach a concentration of some thousands forming unit (CFU) per gram. Studies on French (Tysset and Rousseau, 1981) and Argentinian honeys (Iurlina and Fritz, 2005) showed an average value about 200–250 CFU/g for bacteria and 100–150 CFU/g for fungi. Regarding Italian honeys, lower values of both microbial groups were reported (Piana et al., 1991), even though Farris et al. (1986), which specifically analyzed the bacterial component of Sardinian honeys, detected *Bacillus* spp. at concentrations of approximately 1000 spores/g. So far, the microorganisms detected in honey belong

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to several bacterial (Rozanska, 2011) as well as filamentous fungal (Kačániová et al., 2009) and yeast (Carvalho et al., 2006) species. Other studies carried out on this topic have been mainly forwarded to the hygienic implications and many authors focused on the presence of *Clostridium botulinum* (Saraiva et al., 2012) due to the risk of infant botulism for children below one year old.

Honey is often used as a food ingredient and its microbial load may be transferred to complex matrices where some microorganisms may found the optimal conditions to develop. The knowledge of the microbial composition and the level of the species (and strains), relevant during transformation and/or conservation of the food matrices, may assume a paramount importance for the correct management of the process. Furthermore, the progressive market penetration of foreign honeys, often with a lower quality, increased the interest towards the complete characterization in order to check quality, sanitation and authenticity of the local product.

In light of the above reasons, the quality of honey depends not only on the physical and chemical properties well defined by EC Directive 2001/110, but also on the microbiological aspects largely ignored by the EU legislation. The aim of this study was to deepen the knowledge on the microbial community of untreated nectar honeys and honeydew honeys of different botanical and geographical origin collected in southern Italy.

2. Materials and methods

2.1. Sample collection

A total of 38 artisanal honey (31 nectar honeys and 7 honeydew honeys) samples were purchased from producers, transported to our laboratories and stored in dark conditions at refrigeration temperature until analysis. All honey samples were not apparently characterized by any sign of alteration.

2.2. Microbial counts

Osmophilic microorganisms were counted after homogenization of samples (25 g) in a 30% (w/v) glucose solution (sample/ diluent 1:9) following the indications ISO 21527-2 to avoid shock of cells and to recover sub-lethally injured cells. The first dilution of nectar honeys and honeydew honeys was obtained with a stomacher (BagMixer 400, Interscience, Saint Nom, France) for 2 min at the highest speed. Cell suspensions were spread plated and incubated as follows: total (osmophilic and osmotolerant) yeasts (TY) on tryptone glucose yeast extract agar (TGY), incubated aerobically at 25 °C for 7 d (Beuchat et al., 2001); osmophilic bacteria (OB) on De Whalley Agar (DWA), incubated aerobically at 25 °C for 7 d.

All other microorganisms were recovered by homogenization of samples (25 g) in peptone water. Cell suspensions were plated and incubated as follows: total mesophilic count (TMC) spread on plate count agar (PCA), incubated aerobically at 30 °C for 72 h; filamentous fungi (FF) spread on potato dextrose agar (PDA), incubated aerobically at 25 °C for 21 d; lactic acid bacteria (LAB) poured on glucose M17 (GM17) agar, incubated anaerobically with the AnaeroGen AN25 system at 30 °C for 72 h; *Enterobacteriaceae* poured on violet red bile glucose agar (VRBGA), incubated anaerobically by overlay agar at 37 °C for 24 h; clostridia on reinforced clostridial medium (RCM) by 3 × 3 Most Probable Number (MPN) procedure (FDA BAM, 2006).

Except VRBGA, all media used for bacterial growth were supplemented with cycloeximide (170 ppm) and biphenyl (1 g/L) to inhibit the growth of yeasts and moulds, while all media used for fungal growth were supplemented with chloramphenicol (0.1 g/L) to inhibit bacteria growth. Media were purchased from Oxoid (Basingstoke, UK) and chemicals by Sigma–Aldrich (Milan, Italy). Analyses were performed in duplicate.

2.3. Isolation, grouping and identification of bacteria

After growth, approximately five colonies with the same appearance (colour, morphology, edge, surface and elevation) were collected from count plates of each sample. Presumptive *Enterobacteriaceae* and clostridia were not isolated. The isolates were purified by successive sub-culturing and the purity was checked microscopically. After growth, the cultures were phenotypically characterized by cell morphology, Gram reaction (KOH method) and catalase (determined by transferring fresh colonies from a Petri dish to a glass slide and adding H_2O_2 5%, v/v). Rod, Gram positive, catalase positive bacteria were further characterised for spore formation: cell suspensions were treated at 85 °C for 15 min, (1:10) diluted in Ringer's solution and aliquots of 0.1 ml were spread plated onto Nutrient Agar (NA) (Oxoid) incubated at 32 °C for 48 h.

Otherwise all isolates, excepted Gram-ones, were subjected to biochemical assays and about 40% of the isolates of each pheno-typic group was subjected to molecular analyses.

Genotypic differentiation of selected isolates was first performed by means of randomly amplified polymorphic DNA (RAPD). Genomic DNA for PCR assays was prepared after overnight growth in broth media at 30 °C. Cells were harvested and DNA was extracted by the Instagene Matrix kit (Bio-Rad, Hercules, CA) as described by the manufacturer. Crude cell extracts were used as templates for PCR reactions. RAPD analysis was carried out in a 25-µL reaction mix using primer M13 (Stenlid et al., 1994). Amplifications were performed by means of T1 Thermocycler (Biometra, Göttingen, Germany). PCR products were separated by electrophoresis on 1.5% (w/v) agarose gel (Gibco BRL, Cergy Pontoise, France) and visualized by UV transillumination after staining with SYBR[®] safe DNA gel stain (Molecular probes, Eugene, OR, USA). GeneRuler 100bp Plus DNA ladder (M·Medical Srl, Milan, Italy) was used as a molecular size marker.

All isolates that showed growth onto NA after treatment at 85 °C for 15 min, recognised as spore forming bacteria (SFB), were also analysed by repetitive DNA elements-PCR (rep-PCR) analysis using (GTG)₅ and BOXA1R primer set (Versalovic et al., 1994; Gevers et al., 2001) corresponding to (GTG)₅- and BOX-like elements in bacterial DNA, respectively. All patterns were analyzed using the Gelcompare II software version 6.5 (Applied-Maths, Sin Marten Latem, Belgium). BOXA1R and (GTG)₅ profiles were combined and compared by UPGMA clustering method.

The isolates representative of each cluster were subjected to 16S rRNA gene sequencing. PCR reactions were performed as described by Weisburg et al. (1991). DNA fragments were visualized and the amplicons of about 1600 bp were purified by the QIA-quick purification kit (Quiagen S.p.a., Milan, Italy) and both strands were sequenced using the same primers employed for PCR amplification. DNA sequencing reactions were performed by PRIMM (Milan, Italy). The sequences obtained with forward and reverse primers from each strain were edited and merged into a single sequence covering the entire 16S rRNA gene. In addition, SFB isolates were analysed by restriction fragment length polymorphism (RFLP) of 16S rRNA gene fragment using *Rsa*I, *Cfo*I and *Hinf*I endonucleases (MBI Fermentas, St. Leon-Rot, Germany) (Jeyaram et al., 2011). The sequences were compared by a BLAST search in GenBank/EMBL/DDBJ database.

2.4. Isolation and identification of yeasts

After growth, five colonies per morphology from each sample, or 1-2 isolates for the less numerous groups, were purified onto

the same isolation medium, grouped on the basis of appearance and subjected to genetic characterization.

DNA extraction of all selected isolates was performed as above reported and a first differentiation of yeasts were obtained by RFLP of the region spanning the internal transcribed spacers (ITS1 and ITS2) and the 5.8S rRNA gene. The DNA fragments were amplified with the primer pair ITS1/ITS4 (Esteve-Zarzoso et al., 1999) and subsequently digested with the endonucleases *Cfol*, *Hae*III and *Hinf*I (MBI Fermentas) at 37 °C for 8 h. ITS amplicons as well as their restriction fragments were analysed twice on agarose gel using at first 1.5% (w/v) agarose and then 3% (w/v) agarose in 1× TBE buffer and visualized as above reported. Standard DNA ladders were 1 kb Plus DNA Ladder (Invitrogen) and GeneRuler 50 pb DNA Ladder (MBI Fermentas). One isolate per group was further processed by

sequencing the D1/D2 region of the 26S rRNA gene to confirm the preliminary identification obtained by RFLP analysis. D1/D2 region was amplified with primers NL1 and NL4 (Kurtzman and Robnett, 1998). PCR products visualization and identification was realized as above reported.

2.5. Isolation and identification of filamentous fungi

Fungal colonies were collected and streaked onto PDA, observed at 24-h intervals and once they reached a diameter of approximately 2 mm, transferred to a new Petri dish containing the same substrate. Subsequently, the colonies with different appearance (colour, texture, diffusible pigments, exudates, growth zones, aerial and submerged hyphae, growth rate and topography) were purified

Table 1

Sample Source		Sites of sampling	Media								
code			PCA	GM17	DWA-B	VRBGA	RCM ^a	TGY	DWA-y	PDA	
Miel 01	Er. japonica	Ciaculli (Pa)	$\textbf{7.0} \pm \textbf{2.8}$	$\textbf{3.0}\pm\textbf{0.0}$	_	_	0.36	5420.0 ± 882.0	18600.0 ± 1790.0	_	
Miel 02	Multifloral	Belmonte	3.5 ± 2.1	_	_	_	_	1.0 ± 1.4	_	$\textbf{3.5}\pm\textbf{2.1}$	
		Mezzagno (Pa)									
Miel 03	Multifloral	Naro (Ag)	-	-	-	_	0.36	$\textbf{255.0} \pm \textbf{17.7}$	290.0 ± 45.2	$\textbf{2.5} \pm \textbf{2.1}$	
Miel 04	Citrus spp.	Sciacca (Ag)	-	-	-	_	_	$\textbf{1.0} \pm \textbf{0.0}$	$\textbf{2.0} \pm \textbf{0.0}$	$\textbf{4.0} \pm \textbf{1.4}$	
Miel 05	Multifloral	Trabia (Pa)	$\textbf{2.0} \pm \textbf{0.0}$	1.0 ± 0.0	-	_	0.36	-	-	5.0 ± 1.4	
Miel 06	Multifloral	Trabia (Pa)	$\textbf{4.0} \pm \textbf{0.0}$	-	-	_	-	-	-	$\textbf{27.0} \pm \textbf{4.2}$	
Miel 07	Citrus spp.	Bolognetta (Pa)	47.0 ± 18.0	-	-	-	0.36	476.0 ± 65.1	121.0 ± 29.7	5.5 ± 2.1	
Miel 08	Multifloral	Belmonte Mezzagno (Pa)	$\textbf{6.5} \pm \textbf{2.1}$	-	-	-	-	2.0 ± 1.4	23.0 ± 2.8	$\textbf{3.0} \pm \textbf{0.0}$	
Miel 09	Hed. coronarium	Santa Cristina (Pa)	$\textbf{3.0} \pm \textbf{1.4}$	_	_	_	0.36	5.5 ± 2.1	216.0 ± 55.2	$\textbf{2.0} \pm \textbf{1.4}$	
Miel 10	Citrus spp.	Partinico (Pa)	1.0 ± 1.4	_	_	_	0.36	8.5 ± 2.1	361.0 ± 14.8	5.5 ± 3.5	
Miel 11	R. pseudoacaciae	Caronia (Me)	$\textbf{2.0} \pm \textbf{1.4}$	_	_	_	0.36	_	_	10.5 ± 2.1	
Miel 12	H. honey	Somma Vesuviana (Na)	$\textbf{39.0} \pm \textbf{1.4}$	-	-	_	-	-	-	1.0 ± 0.0	
Miel 13	H. honey	Somma Vesuviana (Na)	54.0 ± 12.7	_	-	-	0.92	$\textbf{6.5}\pm\textbf{2.1}$	$\textbf{6.5}\pm\textbf{2.1}$	_	
Miel 14	H. honey	Castel di Tusa (Me)	5.5 ± 2.1	_	_	_	0.36	_	_	1.0 ± 0.0	
Miel 15	Multifloral	Benevento (Bn)	20.5 ± 3.5	_	_	_	_	6.5 ± 3.5	3.5 ± 2.1	5.0 ± 0.0	
Miel 16	Multifloral	Somma Vesuviana (Na)	9.5 ± 2.1	1.0 ± 1.4	-	-	0.36	17.5 ± 3.5	292.0 ± 61.5	-	
Miel 17	Multifloral	Portici (Na)	7.0 ± 0.0	$\textbf{2.0} \pm \textbf{0.0}$	_	_	_	156.0 ± 22.6	$\textbf{306.0} \pm \textbf{36.8}$	_	
Miel 18	Cas. sativa	Monteforte (Sa)	7.5 ± 2.1	2.0 ± 0.0	_	_	0.36	_	_	_	
Miel 19	H. honey	Somma Vesuviana (Na)	64.0 ± 2.8	13.5 ± 2.1	-	-	0.36	19.0 ± 7.1	550.0 ± 8.5	_	
Miel 20	H. honey	Somma Vesuviana (Na)	30.5 ± 2.1	3.0 ± 0.0	-	-	0.36	$\textbf{63.5} \pm \textbf{23.3}$	288.0 ± 45.3	_	
Miel 21	Euc. camaldulensis	Naro (Ag)	1.0 ± 0.0	_	_	_	_	3.0 ± 0.0	43.0 ± 1.4	_	
Miel 22	Citrus spp.	Sciacca (Ag)	1.0 ± 0.0 1.0 ± 1.4	_	_	_	_	1.0 ± 0.0	_	4.5 ± 2.1	
Miel 23	Car. defloratus	Naro (Ag)	1.0 ± 1.4	_	_	_	0.92	-	_	1.0 ± 1.4	
Miel 24	Multifloral	Naro (Ag)	1.0 ± 0.0	_	_	_	_	_	$\textbf{2.0} \pm \textbf{0.0}$	$\textbf{2.0} \pm \textbf{1.4}$	
Miel 25	Euc. camaldulensis	Lercara Friddi (Pa)	$\textbf{3.0} \pm \textbf{0.0}$	_	_	_	_	$\textbf{782.0} \pm \textbf{31.1}$	874.0 ± 65.1	1.0 ± 1.4	
Miel 26	Car. defloratus	Roccapalumba (Pa)	$\textbf{318.0} \pm \textbf{1.4}$	$\textbf{88.0} \pm \textbf{8.5}$	_	_	_	19.0 ± 5.7	_	_	
Miel 27	H. honey	Finale di Pollina (Pa)	14.0 ± 4.2	1.0 ± 0.0	_	_	_	7.5 ± 2.1	540.0 ± 84.9	5.5 ± 2.1	
Miel 28	Citrus spp.	Santa Flavia (Pa)	108.0 ± 14.1	10.0 ± 2.8	_	2.0 ± 1.4	_	16.5 ± 4.95	56.0 ± 5.7	$\textbf{22.0} \pm \textbf{8.49}$	
Miel 29	Euc. camaldulensis	Piana degli Albanesi (Pa)	9.5 ± 3.5	_	-	_	-	-	-	3.0 ± 1.4	
Miel 30	Multifloral	Castronovo di Sicilia (Pa)	-	_	-	-	-	_	-	1.0 ± 1.4	
Miel 31	Multifloral	Montemaggiore (Pa)	1.0 ± 0.0	_	_	_	_	_	_	_	
Miel 32	Multifloral	Portella di mare (Pa)	1.0 ± 0.0 1.0 ± 0.0	_	_	_	_	_	_	3.0 ± 1.4	
Miel 33	Hed. coronarium	Balata di Baida (Tp)	3.0 ± 0.0	_	_	_	_	_	_	3.0 ± 1.4 3.0 ± 1.4	
Miel 34	Hed. coronarium	Balata di Baida (Tp)	3.0 ± 0.0	_	_	2.5 ± 2.1	0.36	$\textbf{2.0} \pm \textbf{0.0}$	1.0 ± 1.4	3.0 ± 1.4 4.0 ± 1.4	
Miel 35	H. honey	Boscotrecase (Na)	35.5 ± 2.1	6.5 ± 2.1	_	2.5 ± 2.1 -	-	741.0 ± 12.7	1.0 ± 1.4 1600.0 ± 148.0	-	
Miel 36	Multifloral	Boscotrecase (Na)	8.5 ± 3.5	21.5 ± 2.1	_	_	_	_	-	$\textbf{2.0} \pm \textbf{0.0}$	
Miel 37	Multifloral	Balata di Baida (Tp)	-	-	_	_	_	_	_		
Miel 38	Euc. camaldulensis	Balata di Baida (Tp)	_	_	_	_	_	_	_	$\textbf{2.0} \pm \textbf{0.0}$	

Results indicate mean values \pm S.D.

^a As estimated by MPN. Abbreviation: PCA, plate count agar added with cycloeximide and biphenyl for total mesophilic counts; DWA-b, De Whalley Agar added with cycloeximide and biphenyl for osmophilic bacteria counts; GM17, glucose M17 added with cycloeximide and biphenyl for lactic acid bacteria counts; TGY, Tryptone Glucose Yeast extract agar added with chloramphenicol for total (osmophilic and osmotolerant) yeasts; DWA-y, De Whalley Agar added with chloramphenicol for osmophilic yeast counts; PDA, Potato Dextrose Agar added with chloramphenicol for filamentous fungi counts; Me, Messina; Pa, Palermo; Ag, Agrigento; Na, Napoli; Tp, Trapani; Bn, Benevento; *R., Robinia; Car., Cardunculus; Cas., Castanea; Euc., Eucalyptus; H., Honeydew honey; M.,* Multifloral; *Er., Eriobotrya; Hed., Hedysarum.* Symbols: –, under the detection limit.

Table 2

Morphological and biochemical characterization of presumptive LAB found in honeys and honeydew honeys.

Character	Clusters		
	A	В	С
	<i>n</i> = 28	n = 1	<i>n</i> = 10
Morphology	Coccus tetrads	Coccus short chain	Coccus short chain
Mobility	_	-	_
Catalase	_	-	_
Gram reaction	+	+	+
Growth			
15 °C	+	+	+
45 °C	_	_	+
pH 9.2	+	+	+
6.5% NaCl	+	_	+
CO ₂ from glucose	_	_	_
Growth in presence of pentose sugars	_	_	-

Symbols: +, positive; -, negative to the test.

to homogeneity after several sub-culturing steps onto PDA. All colonies were subjected to microscopic analysis (Barnett and Hunter, 1998) through a light microscope (Carl Zeiss Ltd). Five isolates, or less for the less numerous morphological groups, were subjected to genetic analysis. Genomic DNA was extracted from single-spore cultures following a standard cetyl-trimethylammonium-bromide (CTAB) based protocol (O'Donnell et al., 1998). Filamentous fungi were analyzed by RFLP of the region spanning the internal transcribed spacers (ITS1 and ITS2) and the 5.8S rRNA gene. The DNA fragments were amplified with the primer pair ITS1F (Gardes and Bruns, 1993) and ITS4 (White et al., 1990). The amplicons were then digested with the endonucleases CfoI and HaeIII (MBI Fermentas) at 37 °C for 8 h. ITS products, as well as restriction fragments were analyzed on agarose gels using 2% (w/v) agarose in 1× TBE buffer. Gels were treated and visualised as above reported. One isolate per group was processed by sequencing of the 5.8S-ITS rRNA region to confirm the morphological identification.

2.6. Biodiversity indexes

The biodiversity of the overall microbial load was evaluate with Margalef's index of genotypes richness (R), Shannon-Weaver's index of diversity (H') and Simpson's index of dominance (D), calculated as proposed by Ventorino et al. (2007).

3. Results

3.1. Microbial counts

The viable counts of the different microbial groups investigated in this study are reported in Table 1. Although growth of osmophilic bacteria on DWA resulted negative, a positive bacterial development was registered on PCA for the majority of samples. In several samples, only one colony contributed to TMC. However, except sample Miel 36, the highest levels of microorganisms were detected on PCA. Enterobacteriaceae were found only in two samples. The presence of clostridia was found in 15 samples, with the highest values estimated (0.92 MPN/g) for sample Miel 13 and Miel 23. The sample Miel 26 showed the highest bacterial richness. In general, yeasts counted onto DWA were at higher numbers than those registered on TGY, but an opposite observation was made for samples Miel 07, Miel 15, Miel 26 and Miel 34. Several samples resulted positive for the presence of moulds, but only samples Miel 6, Miel 11 and Miel 28, showed concentrations higher than 10 CFU/g. Twelve samples showed the presence of presumptive LAB and their concentrations were in the range 1.0 \pm 0.0–88.0 \pm 8.5 CFU/g. Members of the Enterobacteriaceae family were found only in samples Miel 28 and Miel 34.

3.2. Isolation and identification of bacteria

A total of 464 pure cultures were isolated and purified to homogeneity on the same media used for the plate counts. All cultures were subjected to a preliminary microscopic inspection and separated in three main groups: 423 rod shaped, Gram-positive, catalase positive, spore forming bacteria considered as presumptive Bacillus spp.: two rod shaped. Gram-negative, catalase-positive bacteria: 39 coccus shaped. Gram-positive, catalase negative bacteria considered as presumptive LAB. Due to the limited number of isolates and their different isolation sample, the Gram-negative isolates were subjected to the 16S rRNA gene sequencing without any differentiation at strain level: both isolates were identified as Klebsiella pneumonia (Acc. No. KC692177, KC692181). All other strains were further subjected to biochemical analysis that allowed the subdivision of the presumptive LAB and presumptive Bacillus spp. into three (Table 2) and seven (Table 3) groups, respectively. Furthermore, about 40% of the isolates of each phenotypic group was subjected to RAPD analysis. The resulting dendrograms (Figs. 1 and 2), at a similarity level of 80%, showed that presumptive LAB were divided into three clusters, while presumptive Bacillus spp.

Table 3

Morphological and biochemical characterization of aerobic spore forming bacilli found in honey and honeydew honeys.

Characters	Clusters										
	D (<i>n</i> = 198)	E (<i>n</i> = 50)	F (<i>n</i> = 10)	G(n = 68)	H (<i>n</i> = 95)	I (<i>n</i> = 1)	L (<i>n</i> = 1)				
Cell and spore morphology:											
Vegetative cells	Rod	Rod	Rod	Rod	Rod	Rod	Rod				
Spores	+	+	+	+	+	+	+				
Biochemical tests:											
Catalase	+	+	+	+	+	+	+				
Gram reaction	+	+	+	+	+	+	+				
Anaerobic growth	-	+	+	-	-	+	+				
Nitrate reduction	+	+	+	V	-	+	+				
Voges-Proskauer	+	+	+	-	+	-	+				
Starch hydrolysis	+	+	+	+	-	+	+				
Growth at 50 °C	+	-	+	-	+	-	-				
Growth at 65 °C	-	_	-	_	-	-	_				
Growth in 7% NaCl	+	+	+	+	+	+	_				
Acid and gas in glucose	_	-	_	_	-	-	+				

Symbols: V, variable; +, positive to the test; -, negative to the test.

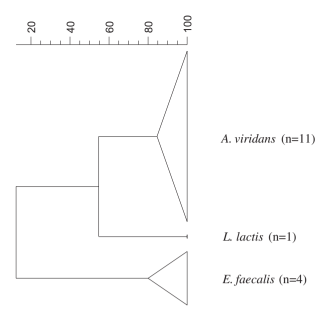


Fig. 1. Dendrogram obtained from RAPD-PCR patterns of presumptive LAB. The values between brackets refer to the number of isolates in each cluster. Upper line indicates the percentage of similarity.

into 33 clusters. The 16S rRNA gene of each presumptive LAB strain was sequenced and the following species were identified: Aerococcus viridans (phenotypic group A, Acc. No. KC692208); Lactococcus lactis (phenotypic group B. Acc. No. KC692209); and Enterococcus faecalis (phenotypic group C, Acc. No. KC692178, KC692183). The presumptive Bacillus spp. strains were further investigated by REP-PCR; (GTG)5 and BOXA1R profiles were combined and the comparison among all isolates recognized 42 different clusters (Table 4). The representative strains of each cluster were analyzed by 16S rRNA gene sequencing (Table 4). The direct comparison performed by BLAST search determined the identification at species level of only five strains: Paenibacillus polymyxa (ML227), Bacillus simplex (ML384), Bacillus pumilus (ML374) and Bacillus licheniformis (ML103A and ML104B). For the other strains indeed also the RFLP analysis was performed and the results were compared with restriction fragments size proposed by Jeyaram et al. (2011). The species Bacillus amyloliquefaciens, Bacillus subtilis, Bacillus cereus, Bacillus thuringiensis, Bacillus licheniformis, Bacillus megaterium and B. pumilus were clearly identified, but the strain ML582, characterized by a 16S rRNA gene sequence homology close to B. amyloliquefaciens/B. subtilis and a RFLP profile similar to B. megaterium, remained unspeciated.

3.3. Isolation and identification of yeasts

Based on morphology, 1027 yeast colonies were collected from TGY and DWA (409 and 618, respectively). Three hundred and twenty-three isolates were then subjected to the molecular identification. After restriction analysis of 5.8S-ITS region, the isolates were clustered in nine groups (Table 5). The isolates of groups III and IX were identified as *Debaryomyces hansenii* and *Zygosacchar-omyces rouxii* by comparison of restriction bands with those available in literature (Esteve-Zarzoso et al., 1999; Carvalho et al., 2006). The other groups could not be identified by RFLP analysis and the identification at species level was concluded by sequencing of D1/ D2 domain of the 26S rRNA gene which recognised three species: *Zygosaccharomyces mellis* for the isolates included into the groups IV, V, VI, VII and VIII; *Aureobasidium pullulans* for group I; *Crypto-coccus uzbekistanensis* for group II. *Z. mellis* (group IV) was the most isolated species and it represented the majority of yeast population in the great part of samples. On the contrary, the other species were less numerous and they were detected only from six different honeys. A total of 23 samples were positive for yeast presence and the samples Miel 19 and Miel 35, both honeydew honeys, showed the highest yeast diversity at strain level.

3.4. Isolation and identification of filamentous fungi

A total of 117 filamentous fungi were collected and divided into 17 groups after microscopic inspection (Table 6). The results of the restriction analysis of 5.8S-ITS region using the endonucleases CfoI and HaeIII confirmed the subdivision of the filamentous fungi into 17 groups. One isolate per group was subjected to the sequencing of the 5.8S-ITS rRNA gene that clearly identified the species Alternaria alternata (group I), Aspergillus niger (group III), Aspergillus proliferans (group IV), Aspergillus spelunceus (group V), Chaetomium globosum (group VI), Cladosporium cladosporioides (group VII), Daldinia concentrica (group VIII), Emericella discophora (group IX), Emericella qinqixianii (group X), Penicillium corylophilum (group XII), Penicillium decumbens (group XIII), Penicillium italicum (group XIV), Penicillium polonicum (group XVI) and Penicillium echinulatum (group XVII). The isolates of groups II, XI and XV were identified at genus level as Arthrinium, Emericella and Penicillium, respectively. The species most frequently isolated were Pen. corylophilum and Asp. niger. The samples Miel 7 (Citrus spp.) and Miel 3, Miel 5 and Miel 15 (multifloral) were the richest sources of fungal diversity among the 26 samples positive to fungal isolation.

3.5. Distribution of strains and biodiversity indexes

The distribution of the microorganisms isolated from honeys is reported in Table 7. *B. amyloliquefaciens*, *B. megaterium*, *B. pumilus*, *Aspergillus niger*, *Penicillium corylophilum* and *Z. mellis* were commonly found in several samples of different botanical origin. Margalef's, Shannon-Weaver's and Simpson's indexes (Table 8) showed the highest microbial diversity for multifloral, *Citrus* and *He. coronarium* honeys. In particular, the highest R and H' and the lowest D values were registered for the multifloral honeys.

4. Discussion

Honey retains a natural image and the increasing trend in consumption could be also attributed to a new life style with people demanding more natural foods for the beneficial effects of these products (Arvanitoyannis and Krystallis, 2006). However, the quality of honey could be compromised by the hygienic practices during harvest and extraction as well as time and conditions of storage (Snowdon and Cliver, 1996). The intrinsic properties of honey affect the growth and survival of several microorganisms; thus, they influence the species proportions, but some microbial species, especially bacteria with pathogenic characteristics, have been isolated from honey samples. Hence, the interest towards the microbial composition of commercial honeys is of paramount importance, even though it is considered a food ingredient and it is consumed in low amounts. In order to better examine the living microbial communities associated with honey, 38 samples were collected in different cities of southern Italy and analysed by culture-dependent methods. This because the presence of undesired microorganisms that are able to develop colonies undoubtedly provide a clear indication of the hygienic safety of honey.

Despite the stressing conditions of honey, the microbiological analyses revealed the presence of several microbial groups. The results of our study, conducted on honey and honeydew honey samples, showed that 33 of them hosted low loads of bacteria, with

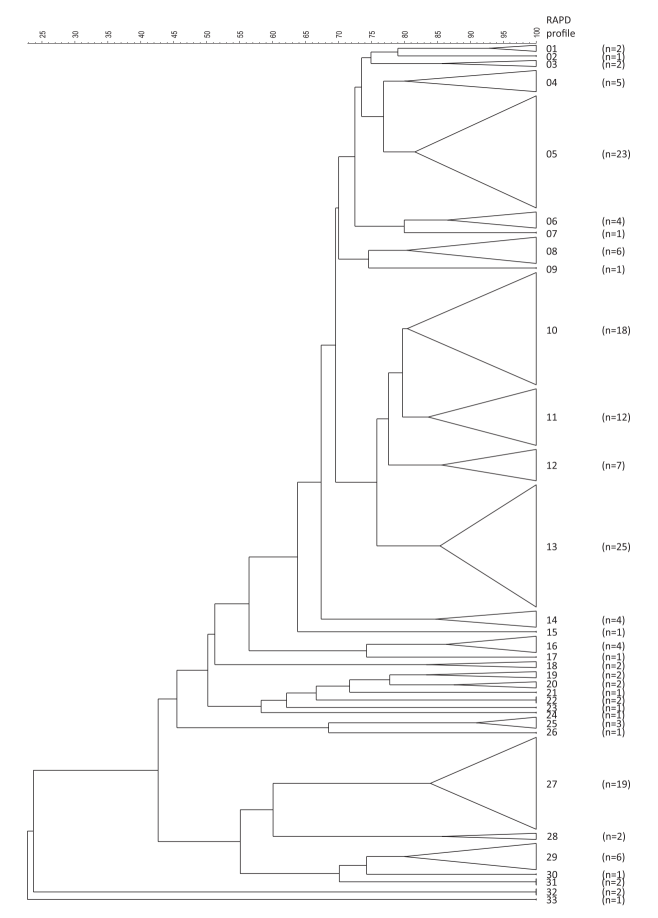


Fig. 2. Dendrogram obtained from RAPD-PCR patterns of presumptive *Bacillus* spp. The values between brackets refer to the number of isolates in each cluster. Upper line indicates the percentage of similarity.

Table 4

Molecular identification of spore forming bacteria.

Strain	Phenotypic cluster	RAPD cluster	GTG/BOXA1R cluster	Identification by 16S and BLAST search	Final identification by RFLP	Acc. No.	No. of isolate
ML101A	E	29	35	B. cereus/B. thuringiensis	B. cereus	KC692164	2
ML101B	D	25	20	B. subtilis/B. mojavensis	B. subtilis	KC692170	3
ML102B	D	28	41	B. subtilis/B. mojavensis	B. subtilis	KC692194	2
ML103A	F	32	33	B. licheniformis		KC692195	2
ML104B	F	18	1	B. licheniformis		KC692185	2
ML105	Н	23	19	B. pumilus/B. safensis	B. pumilus	KC692196	1
ML106A	D	33	10	B. subtilis/B. amyloliquefaciens	B. amyloliquefaciens	KC692197	1
ML106B	D	19	28	B. subtilis/B. amyloliquefaciens	B. amyloliquefaciens	KC692198	2
ML208	Е	29	35	B. cereus/B. thuringiensis	B. cereus	KC692193	4
ML211	Н	5	23	B. pumilus/B. altitudinis	B. pumilus	KC692172	5
ML215	D	1	27	B. subtilis/B. mojavensis	B. subtilis	KC692182	2
ML227	L	26	34	P. polymyxa		KC692186	1
ML233	Е	5	24	B. thuringiensis/B. cereus	B. thuringiensis	KC692184	6
ML235	G	31	11	B. megaterium/B. aryabhattai	B. megaterium	KC692187	2
ML252	Н	5	25	B. pumilus/B. altitudinis	B. pumilus	KC692165	3
ML254	E	30	36	B. cereus/B. thuringiensis	B. cereus	KC692199	1
ML256	Н	5	25	B. pumilus/B. altitudinis	B. pumilus	KC692162	6
ML257	G	20	12	B. megaterium/B. aryabhattai	B. megaterium	KC692200	1
ML258	G	22	14	B. megaterium/B. aryabhattai	B. megaterium	KC692166	2
ML259	E	6	37	B. cereus/B. thuringiensis	B. cereus	KC692188	4
ML265	D	17	3	B. subtilis/B. amyloliquefaciens	B. amyloliquefaciens	KC692168	1
ML267	Е	5	39	B. cereus/B. thuringiensis	B. cereus	KC692161	2
ML270	Н	5	32	B. pumilus/B. safensis	B. pumilus	KC692158	1
ML272	E	15	21	B. cereus/B. thuringiensis	B. cereus	KC692201	1
ML274	D	11	6	B. subtilis/B. amyloliquefaciens	B. amyloliquefaciens	KC692159	11
ML275	G	21	13	B. megaterium/B. aryabhattai	B. megaterium	KC692202	1
ML323	D	16	7	B. amyloliquefaciens/B. subtilis	B. amyloliquefaciens	KC692203	4
ML345	D	10	3	B. amyloliquefaciens/B. subtilis	B. amyloliquefaciens	KC692167	2
ML353	Н	4	22	B. pumilus/B. altitudinis	B. pumilus	KC692160	5
ML361	D	13	5	B. subtilis/B. amyloliquefaciens	B. amyloliquefaciens	KC692163	25
ML374	Н	3	30	B. pumilus	5 1 5	KC692204	2
ML376	D	10	40	B. amyloliquefaciens/B. subtilis	B. amyloliquefaciens	KC692189	2
ML384	Ι	11	26	B. simplex	5 1 5	KC692191	1
ML390	D	9	4	B. subtilis/B. amyloliquefaciens	B. amyloliquefaciens	KC692192	1
ML451	D	10	2	B. amyloliquefaciens/B. subtilis	B. amyloliquefaciens	KC692171	5
ML456	D	10	8	B. subtilis/B. amyloliquefaciens	B. amyloliquefaciens	KC692174	2
ML471	D	2	42	B. amyloliquefaciens/B. subtilis	B. amyloliquefaciens	KC692205	1
ML474	D	14	16	B. subtilis/B. amyloliquefaciens	B. subtilis	KC692190	4
ML477	H	8	17	B. pumilus/B. safensis	B. pumilus	KC692169	2
ML479	G	20	9	B. megaterium/B. aryabhattai	B. megaterium	KC692206	1
ML482	G	7	15	B. megaterium/B. aryabhattai	B. megaterium	KC692173	1
ML484	H	12	31	B. pumilus/B. altitudinis	B. pumilus	KC692175	7
ML568	H	8	29	B. pumilus/B. altitudinis	B. pumilus	KC692176	4
ML581	D	10	3	B. subtilis/B. amyloliquefaciens	B. amyloliquefaciens	KC692179	7
ML582	D	27	38	B. amyloliquefaciens/B. subtilis	Not resolved	KC692180	19
ML618	H	24	18	B. pumilus/B. safensis	B. pumilus	KC692207	15

318 CFU/g as maximum level of concentration, a value close to that reported by other researchers (Gomes et al., 2011). Presumptive LAB were detected in twelve honeys and only two samples resulted positive for the presence of *Enterobacteriaceae*. Surprisingly, any bacterial growth was not detected on DWA medium, generally employed to reveal the osmophilic bacteria, probably because the composition of this medium is still too stressing to allow cell division. Clostridia, even though at low levels, were observed in about 40% of the samples analysed. Some works conducted on the search of *Clostridium* spp. reported negative results (lurlina and Fritz, 2005; Gomes et al., 2011). However, some Italian honeys have been found to host SFB at levels comparable to those of our study (Piana et al., 1991), but higher concentrations, till two orders of magnitude, have been also reported for sugar rich matrix (Finola et al., 2007).

Except three samples, all other honeys and honeydew honeys were found positive for the presence of yeasts or moulds, with maximum concentrations of 18,600 and 27 CFU/g, respectively. The low value obtained for moulds confirmed that the growth of these organisms is limited in honey. The maximum yeast load was

registered for the *Er. japonica* honey. This finding is not surprising because it is a product harvested in winter when the moisture content can be higher due to hygroscopic properties of honey (Zumla and Lulat, 1989). The other samples showed fungal levels closer to those found by other authors (Piana et al., 1991; Iurlina and Fritz, 2005).

All bacteria isolated were divided into three main groups after molecular identification. The Gram-negative isolates were identified as *Klebsiella pneumoniae*. LAB were grouped into three groups including one species each (*A. viridans, L. lactis* and *E. faecalis*). *K. pneumoniae* and *L. lactis* have been isolated from the digestive tract of *Apis cerana* (Nada et al., 2010; Ahn et al., 2012), but even though it is different from *A. mellifera*, it might be reasonably supposed that the same species may be associated to *A. mellifera* and, consequently, contaminate honey products. To our knowledge, no previous studies reported the presence of these species in honeydew honey and this is also the first finding of *K. pneumoniae*, *L. lactis, E. faecalis* and *A. viridans* from nectar honeys.

Seven groups of SFB were obtained and they included several species within *Bacillus* genus and *P. polymyxa*. *B. amyloliquefaciens*

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IGY NO. OF Isolates		101 Miel (2.4)) 7, 8, 9, 15, 16, 17, 22, 25, 26, 27, 28)) 7, 8, 9, 15, 16, 17, 22, 25, 26, 27, 28)
				s	
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	300 + 175 + 125 Aureobasidium pullulans	300 + 175 + 125 350 + 270	300 + 175 + 125 350 + 270 325 + 325	300 + 175 + 125 350 + 270 325 + 325 400 + 260 + 190	300 + 175 + 125Aureobasidium350 + 270cryptococcus350 + 270cryptococcus325 + 325Debaryomyces400 + 260 + 190Zygosaccharomyces530 + 260 + 130Zygosaccharomycesmellismellis
	150 + 150	450 + 150 550 + 70	450 + 150 550 + 70 420 + 150 + 90	450 + 150 550 + 70 420 + 150 + 90 590 + 200 + 60	450 + 150 550 + 70 420 + 150 + 90 590 + 200 + 60 700 + 160 + 60
	200 + 180 + 110 + 60 450 + 150	200 + 180 + 110 + 60 45 320 + 300 55	180 + 110 + 60 300 300 + 50	$ 180 + 110 + 60 \\ 300 \\ 300 + 50 \\ 240 + 190 + 110 $	180 + 110 + 60 300 300 + 50 240 + 190 + 110 390 + 140
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No. of isolates

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Miel (1)

114

Miel (3, 4, 7, 8, 9, 10, 16, 17, 19, 20, 21, 24, 25, 27, 28, 34, 35)

resulted the species with the highest number of isolates. Several studies reported the isolation of Bacillus species in honeys (Iurlina and Fritz, 2005; Alippi and Reynaldi, 2006) but this study represents the first report on the identification of Bacillus spp. in honeydew honeys. With the exception of two species (Bacillus anthracis and many *B. cereus* toxin-producer strains). *Bacillus* group is considered safe. So far, honey is not known to have been involved in disease caused by B. cereus, but López and Alippi (2010) considered honeys contaminated with B. cereus and B. megaterium as possible vectors of foodborne illnesses. However, thanks to the ability of several Bacillus strains to produce antibiotics, bacteriocins, or antifungal compounds, they find application to pursue agricultural and healthcare purposes (Alfonzo et al., 2012; Duc et al., 2004). Some strains of P. polymyxa (Lee et al., 2009), B. subtilis (Sabatè et al., 2009), B. cereus, B. licheniformis and B. megaterium (Alippi and Reynaldi, 2006), isolated from honey samples or different apiarian sources, were successfully tested against Paenibacillus larvae. B. amyloliquefaciens is considered able to inhibit Ascosphaera apis (Brittes Benitez et al., 2012). P. larvae and A. apis are two common pathogens causing American foulbrood and chalkbrood in honey bee colonies (Sabatè et al., 2009). Today diseases and parasites are considered one of the main reasons of the decrease in bee population (Genersch, 2010) and the selection of microbial strains with antagonistic effect represents an important strategy for the biological control of these dangerous organisms (Yoshiyama et al., 2013).

Nine veast species were identified, but only Z. rouxii and D. hansenii were directly identified by 5.8S rRNA gene RFLP: for the other species the sequencing of the D1/D2 domain of the 26S rRNA gene was necessary, because atypical restriction profiles of 5.8S-ITS were registered. An atypical polymorphism for this region is not surprising, since many authors observed this behaviour in several yeasts (Solieri et al., 2007; Tofalo et al., 2009). The two osmotolerant Z. mellis and Z. rouxii are commonly associated with the honey environment (Deak, 2007; Carvalho et al., 2006), but no previous detection of A. pullulans, C. uzbekistanensis and D. hansenii in honey sources are available in literature. Z. mellis is a spoiling agent of high sugar foods and honeys (Wrent et al., 2010). This species is not able to use sucrose for growth (Kurtzman and Fell, 1998) and can easily survive in nectar and honey that are rich in glucose and fructose (Tysset and Rautlin de la Roy, 1974).

Filamentous fungi, on the basis of macroscopic, microscopic and molecular analysis were divided into 17 groups consisting of eight different genera (Alternaria, Arthrinium, Aspergillus, Chaetonium, Cladosporium, Daldinia, Penicillium and Emericella) with the prevalence of the species Pe. corylophilum and As. niger (50% and 32% of the samples, respectively). Penicillium, Cladosporium, Alternaria and Aspergillus genera are considered common contaminants of honey (Nasser, 2004; Kačániová et al., 2009), while the other species have not been reported vet. In particular, with the exception of Da. concentrica, a wood saprophyte fungus (Boddy et al., 1985), the other species belong to genera known as fungal allergens and mycotoxin producers (Griessler et al., 2010; Moss, 2002).

In conclusion, 35 microbial species associated with honeys and honeydew honeys were identified in this study, confirming that the stressing conditions of honey are highly selective. Thus, this matrix can be considered as a source of microorganisms useful to act in suboptimal conditions, e.g. to carry out the transformation of wastes from the food industry characterized by high concentration of sugars (e.g. molasses) in order to reduce the volume of untreated waste bulks and to valorise these substrates to obtain microbial metabolites with different applications.

The biodiversity indexes revealed some differences among samples within the same botanical origin. The microbiota of multifloral honeys showed the highest values for genotype richness

2 6 18

Miel (13, 35) Miel (1)

4 26 10

Miel (13, 16, 19, 20, 34, 35)

KC692235 KC692232 KC692234

Miel (10) Miel (1)

Zygosaccharomyces mellis Zygosaccharomyces mellis Zygosaccharomyces rouxii

 $\begin{array}{c} 400+250+170\\ 510+260+130\\ 350+260+140 \end{array}$

 $\begin{array}{r} 670 \,+\, 170 \,+\, 60 \\ 400 \,+\, 210 \,+\, 90 \end{array}$

 $\begin{array}{c} 300+220+190+110\\ 390+360+140\\ 290+200+170+90 \end{array}$

820 900 750

ML343 ML534 ML119

III V

 \cong

820

restriction fragments are given in bp.

All values for the 5.8S-ITS PCR and restriction fragments are gi The number reported between brackets refers to sample code.

Viel (28) Miel (19)

Table 6
Molecular identification of filamentous fungi.

RFLP	Isolate	5.8S-ITS	'S Size of restriction fragments		Specie (% identity ^a)	Acc. No.	Source of isolation	No. of
	code PCR		CfoI	HaeIII				isolates
I	ML356	620	360 + 140 + 120	470 + 150	Alternaria alternata (99)	KC692221	Miel 15	1
II	ML193	630	360 + 270	470 + 160	Arthrinium spp. (99)	KC692225	Miel (3, 4, 5, 7, 15)	9
III	ML168B	620	200 + 180 + 130 + 90	300 + 100 + 70 + 50	Aspergillus niger (99)	KC692215	Miel (3, 4, 5, 7, 8, 22, 24, 27, 29,32, 34, 38)	35
IV	ML280	620	330 + 270	290 + 170 + 85	Aspergillus proliferans (100)	KC692212	Miel 7	1
V	ML442	620	210 + 200 + 80 + 50	510 + 100	Aspergillus spelunceus (99)	KC692218	Miel 23	1
VI	ML176	620	280 + 190 + 150	250 + 130 + 80 + 60	Chaetomium globosum (99)	KC692213	Miel 2	1
VII	ML370	600	340 + 260	600	Cladosporium cladosporioides (99)	KC692219	Miel 11	1
VIII	ML286	600	180 + 160 + 130 + 80	600	Daldinia concentrica (99)	KC692211	Miel 8	1
IX	ML297	600	260 + 130 + 80	400 + 100 + 70	Emericella discophora (99)	KC692226	Miel 9	1
х	ML514	600	260 + 190 + 80	430 + 110 + 80	Emericella qinqixianii (99)	KC692210	Miel 33	1
XI	ML488	620	250 + 130 + 90 + 60	410 + 100 + 70	Emericella spp. (96)	KC692217	Miel 25	1
XII	ML369	630	180 + 90	260 + 90 + 70 + 55	Penicillium corylophilum (99)	KC692220	Miel (2, 3, 4,5, 6, 7, 9, 10, 11,12, 14, 22, 28, 29, 30, 32,33, 34, 36)	39
XIII	ML155	620	180 + 90	260 + 90 + 80 + 55	Penicillium decumbens (100)	KC692216	Miel (3, 4)	11
XIV	ML332	620	190 + 100	260 + 95 + 70	Penicillium italicum (99)	KC692222	Miel (10, 15)	3
XV	ML172	630	220 + 180 + 170 + 60	390 + 150 + 90 + 50	Penicillium spp. (95)	KC692214	Miel 5	1
XVI	ML329	630	180 + 170 + 140 + 90	260 + 95 + 70 + 50	Penicillium polonicum (99)	KC692223	Miel (2, 5, 6, 10, 11, 15)	9
XVII	ML291	620	170 + 90	250 + 100 + 70	Penicillium echinulatum (99)	KC692224	Miel 7	1

The number reported between brackets refers to the number code of samples. ^a According to BlastN search of ITS1-5.8S-ITS2 rRNA gene sequences in NCBI database.

Table 7

Distribution of the microorganisms throughout honeys and honeydew honeys of different botanical origin.

Microorganisms	Species									
	Car. defloratus	Cas. sativa	Citrus spp.	Euc .camaldulensis	Er. japonica	H. honey	Hed. coronarium	Multifloral	R. pseudoacacia	
Bacteria:										
Bacillus amyloliquefaciens	х	х	х	х	х	х	х	х		
Bacillus cereus	х				х	х		х		
Bacillus licheniformis					х			х		
Bacillus megaterium	х		х	х		х	х	х		
Bacillus pumilus	х	х		х	х	х		х		
Bacillus simplex		х								
Bacillus subtilis		х			х		х	х		
Bacillus thuringiensis						х		х	х	
Paenibacillus polymyxa								х		
Aerococcus viridans	х									
Lactococcus lactis							х			
Enterococcus faecalis			х			х		х		
Klebsiella pneumoniae			x				х			
Filamentous fungi:										
Alternaria alternata								х		
Arthrinium spp.			х					x		
Aspergillus niger			x	х		х	х	x		
Aspergillus proliferans			x	A		A	A	А		
Aspergillus spelunceus	х		A							
Chaetomium globosum	А							х		
Cladosporium cladosporioides								A	х	
Daldinia concentrica								х	A	
Emericella discophora							х	л		
Emericella gingixianii							x			
Emericella spp.				х			~			
Penicillium corylophilum			x	x		х	х	x	х	
Penicillium italicum			x	A		Λ	~	x	A	
Penicillium polonicum			x					x	х	
Penicillium spp.			^					x	Λ	
Penicillium decumbens			х					x		
Penicillium echinulatum			x x					Λ		
Yeasts:			^							
Aureobasidium pullulans								x		
Cryptococcus uzbekistanensis			v					Λ		
Debaryomyces hansenii	Y		x		Y					
Zygosaccharomyces mellis	x		x	v	х	v	v	v		
	х		х	х	v	х	х	х		
Zygosaccharomyces rouxii					х					

Table 8 Biodiversity indexes

Botanical source	Indexes		
	R	H′	D
Car. defloratus	2.15	1.72	0.24
Cas. sativa	1.67	1.33	0.28
Citrus spp.	2.98	1.97	0.22
E. camaldulensis	1.69	1.37	0.34
Er. japonica	1.67	1.26	0.43
H. honey	1.74	1.70	0.24
He. coronarium	2.73	1.87	0.22
Multifloral	3.91	2.35	0.15
R. pseudoacaciae	1.36	1.21	0.33

Indexes: R, Margalef's index of genotypes richness; H', Shannon-Weaver's index of diversity; D, Simpson's index of dominance.

and diversity indexes, but contemporarily they were characterised by the lowest value of dominance. Some species were detected in almost all honeys and some strains, regardless on the origin of the product, were recognised more than once highlighting their strong adaptation to this matrix. For the species found occasionally, a correlation with the botanical or geographical origin cannot be found, but only an environmental contamination may be supposed. Furthermore, our findings are close to that reported by Snowdon and Cliver (1996) for industrial honey, probably thanks to the correct handling techniques applied by the beekeepers. Nevertheless, the presence of spores of potentially dangerous microorganisms evidenced the need to further study the microbial communities of honey for the hygienic safety of this product. Works are being prepared to evaluate the presence of viable but not cultivable populations in order to provide a more complete view of the microbial complexity of honeys.

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