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Analysis of a Pool of Small Plasmids from Soil Heterotrophic Cultivable Bacterial Communities

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Abstract: In this work the analysis of the plasmid presence on soil aerobic cultivable heterotrophic bacterial communities was carried out checking a panel of 1,200 isolates, in order to establish the frequency of plasmid presence as well as the degree of plasmid flow between strains affiliated to the same or different taxon. Bacterial communities were isolated from two different sites of a 13-year experimental field with a clay-silt texture. Plasmid molecules were detected at low frequency (27 isolates, 2%) with a size ranging between 2 Kb and 40 Kb. The RAPD analysis performed on the plasmid-harboring isolates and the phylogenetic analysis of the whole community using the 16S rRNA gene sequences revealed the existence of transfer of the same plasmids between strains belonging to the same species and, in some cases, to different species of the same genus. As it might be expected, even though the viable cells title did not differ significantly between the two samplings, the overall data disclosed an uneven distribution of both species and plasmid-harboring strains.

Keywords: Horizontal gene transfer, plasmids, r-K strategy, soil.

INTRODUCTION

Soil is a complex and dynamic ecosystem whose functionality is related to the equilibrium existing between chemical, physical, biological parameters and the resident microbial communities. The biodiversity of these communities may undergo fluctuations as a consequence of environmental changes. It has been recognized that one of the key factors responsible for the biodiversity of soil microbial communities and, in general, for microbial evolution is represented by mobile genetic elements (MGE: bacteriophages, transposons and plasmids) that are involved in the horizontal transfer of genetic information (HGT) [1]. In fact, "while point mutations contribute to microbial adaptation, horizontal dissemination of genes has proven to be critical in promoting rapid genomic flexibility and microbial evolution" [2]. Particularly interesting from this viewpoint are plasmids, for the essential role they play in the ecological adaptation of Bacteria and Archaea; indeed, they can contribute to shape prokaryotic genomes, "promoting intra- and inter-species variability and distributing functional genetic modules" [3].

Genomics approaches allowed to disclose a large and untapped diversity of plasmids inhabiting plant-associated or soil bacteria. "Surveys on the presence of plasmids from soil and plant-associated bacteria have been performed and revealed that a considerable portion of bacteria from different environments carried plasmids; for examples, approximately 18% of bacterial isolates from the phytosphere of sugar beets

were found to contain plasmids" [4]. Furthermore, although the function of most of plasmid-borne genes is still unknown [5], there is a general agreement that MGE add some, often small or even not measurable, metabolic burden to their host. "Traits conferring an improved fitness or ability to colonize environmental niches are often located on conjugative MGE; consequently, the prevalence of plasmids indicates that they can benefit bacteria in the environment" [6].

Rhizosphere, together with soil, is one of the main "hot" spots for gene transfer activity performed by bacteria. This is due to different factors, including the enhanced nutrient input and water fluxes that might stimulate bacterial metabolic activities. The enhanced conjugative transfer of chromosomal genes between *Pseudomonas* spp. in the wheat rhizosphere in respect to bulk soil was previously reported [7]. Another example of bacterial metabolic abilities, which very likely have been rather recently evolved and spread through HGT is the capability to perform biodegradation of man-made xenobiotic compounds [8].

The plasmids ecology is still poorly understood and we know little of their distribution and diversity. In spite of the importance of plasmid molecules, an extensive analysis of the presence and frequency of plasmids in large natural cultivable microbial communities has not been performed up to now. Therefore, the aim of this work was to analyze the presence of plasmids in a large heterotrophic cultivable bacterial community isolated from soil and to check the degree of genetic flow between strains belonging to the same or different species/genus. To this purpose we used the experimental strategy schematically represented in Fig. (1).

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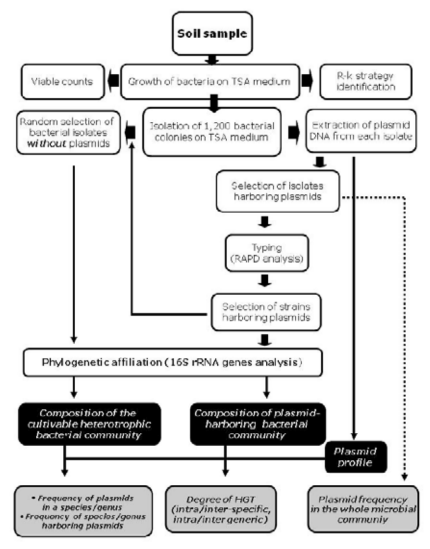


Fig. (1). Schematic representation of the overall experimental strategy used in this work.

MATERIALS AND METHODS

Soil Sampling and Processing

Bacteria analyzed in this work (Table 1) were isolated from six soil samples collected from the top and the bottom of a 13-years experimental field located in the coastal hills of Marche (Agugliano, Ancona, Central Italy). The six soil samples were referred to as 2.2 A, 2.2 B, 2.2 C and 4.1 A, 4.1 B and 4.1 C (from the top and the bottom of the experimental field, respectively). The Agugliano soil is a Calcaric Glevic Cambisol with 20% slope [9], which in the first 30 cm has an Ap horizon (that is the homogeneous layer due to plowing) and a clay-silt texture [10]. The soil is managed under a Triticum durum (in winter) and Zea mays (in summer) rotation. Six soil samples were collected on 11 June 2007, during the maize rotation, in no-tillage (NT) system (sod seeding with chemical desiccation and chopping) and unfertilized (UF) soil (0 Kg N ha⁻¹) at 0-20 cm depth. Each soil sample consisted of five soil cores taken inside two NT-UF blocks (top and bottom) of experimental field free from roots and then pooled together. Soil samples were sieved immediately at 2-mm mesh size, kept at 4°C and processed for further analysis within 24 h from sampling.

Culturing of Fast- and Slow-growing Culturable Bacteria

About 1-g (wet weight) of each soil sample was suspended in 10 ml of sterile phosphate-buffered saline (PBS, pH 7.3), homogenized at low speed 3 (Ultra-Turrax Thyristor Regle 50. Janke & Kunkel IKA-Labortechnik) and vortexed for 30 seconds. Then, each sample was transferred into a sterile 100-ml Erlenmeyer flask containing 10 g of glass beads (average diameter, 2 mm previously autoclaved for 20 min at 121°C) and shaken for 1 h at 120 rpm and 28°C to disperse bacteria. The flasks and glass beads were autoclaved for 20 min at 121°C before use. The resulting soil suspension was removed and transferred to a sterile 15-ml Falcon tube. Serial dilutions of this suspension were performed with sterile saline solution (9 g l⁻¹NaCl) from 10⁻¹up to 10⁻⁷. Then, 100 µl aliquots of serially diluted soil suspensions were plated in triplicate on 0.1 tryptic soy broth (TSB, Difco) containing 15 g l⁻¹ agar (0.1 TSA) and 100 µg ml⁻¹ cycloheximide (Sigma) to inhibit fungal growth. Plates were incubated at 28 °C for 6 days. Total culturable bacteria were enumerated on the basis of the r/K strategy concept [11] at day 1, 2 and 6; in this way, three counts per plate were performed, corresponding to three classes (1, 2, and 3) with different

Table 1. List of bacterial isolates analysed in this work; the phylogenetic affiliation of each isolate is also reported along with the 16S rRNA gene accession number.

Isolate	Sampling	Plasmid presence	16S rDNA accession number	First Blast Hit	Phylogenetic affiliation	
1		-	GU808446	AF131549 Streptomyces sp. IM-7082	Streptomyces	
2		-	GU646899	JF700422 Stenotrophomonas sp. WR49	Stenotrophomonas	
9		-	GU831905	FR682931 Stenotrophomonas sp. R-41388	Stenotrophomonas	
16		-	GU814021	JF700422 Stenotrophomonas sp. WR49	Stenotrophomonas	
56		-	GU814032	FJ006929 Microbacterium sp. WPCB194	Microbacterium	
57		-	GU808432	EU366363 Bacillus pumilus strain I5	Bacillus	
85		- GU814025 JF700422 Stenotrophomonas sp. WR49		Stenotrophomonas		
98	224	-	GU808428	JF521654 Rhodococcus sp. CS1	Rhodococcus	
99	2.2.A	-	GU814026	JF700422 Stenotrophomonas sp. WR49	Stenotrophomonas	
116		-	GU831909	JF825992 Bacillus sp. DP5 (2011)	Bacillus	
120		-	GU831884	GU361112 Klebsiella oxytoca strain SHD-1	Klebsiella	
154		-	GU831895	FM162997 Microbacteriaceae bacterium ACEMC 25-3	Microbacterium	
171		-	GU831910	HQ257249 Bacillus sp. SG3	Bacillus	
184		-	GU831890	HQ317157 Paenibacillus polymyxa strain DYJL14	Paenibacillus	
202		-	GU814020	HQ185398 Stenotrophomonas maltophilia strain 5517	Stenotrophomonas	
209		-	GU831908	HQ877451 Stenotrophomonas sp. 33T	Stenotrophomonas	
210		+	GU646895	GU646895 Pseudomonas geniculata	Pseudomonas	
211		-	GU814019	GQ381282 Stenotrophomonas sp. TC7	Stenotrophomonas	
219		- GU646916 HQ406755 Acinetobacter sp. TY14McD		Acinetobacter		
220		- GU814034 GQ369018 Microbacterium sp. T0-YC6750		Microbacterium		
227		-	- GU646917 HQ647282 S. maltophilia strain TS51		Stenotrophomonas	
230		+	GU646884	FJ263916 Acinetobacter johnsonii strain BA2	Acinetobacter	
238		-	GU831885	EU445236 Agrobacterium tumefaciens isolate EFLRI 54	Agrobacterium	
253	2.2.B	+	GU646885	FJ263916 Acinetobacter johnsonii strain BA2	Acinetobacter	
266		-	GU831891	EF120473 Enterobacter cloacae	Enterobacter	
300		-	GU814027	FM207522 Acinetobacter calcoaceticus strain CWS20	Acinetobacter	
308 bis		+	GU646904	FN393790 Acinetobacter lwoffii strain ES-117-3	Acinetobacter	
313		-	GU814024	HQ877451 Stenotrophomonas sp. 33T	Stenotrophomonas	
327		+	GU646886	FJ263916 Acinetobacter johnsonii strain BA2	Acinetobacter	
330		-	GU646918	AY366481 Acinetobacter sp. C1010	Acinetobacter	
343		-	GU831896	EU855207 Enterobacter sp. CTSP29	Enterobacter	
363	†	+	GU646906	AB461770 Enterobacter sp. M429	Enterobacter	

(Table 1) contd....

Isolate	Sampling	Plasmid presence	16S rDNA accession number	First Blast Hit	Phylogenetic affiliation	
385		-	GU831907	HQ877451 Stenotrophomonas sp. 33T	Stenotrophomonas	
386		-	GU831906	HQ877451 Stenotrophomonas sp. 33T	Stenotrophomonas	
397		+	HM046411	FJ263916 Acinetobacter johnsonii strain BA2	Acinetobacter	
408		-	GU814036	EU714376 Microbacterium foliorum strain 720	Microbacterium	
412		-	GU814023	AY599705 Stenotrophomonas sp. TB4-3-II	Stenotrophomonas	
440		-	GU814035	DQ530139 Microbacterium sp. RI48	Microbacterium	
449		-	- GU808436 JN009619 Enterobacter sp. lb11		Enterobacter	
450		-	GU808435	EU999992 Enterobacter ludwigii strain T4384	Enterobacter	
457		+	GU646898	EU999992 Enterobacter ludwigii strain T4384	Enterobacter	
458		-	GU808431	JN036709 Bacillus thuringiensis strain AIMST KBT9-X	Bacillus	
470	+ GU6		GU646887	HQ185399 Stenotrophomonas maltophilia strain 2681	Stenotrophomonas	
479		+ GU646888 HQ694446 Pantoea agglomerans strain AIMST 4.P1.4		Pantoea		
486		+	GU646905	FR774919 Acinetobacter sp. R-45867	Acinetobacter	
489		-	GU814028	GU814028 AB619594 Acinetobacter sp. NCCP 233		
500		+	+ GU646900 EU999992 Enterobacter ludwi		Enterobacter	
505		-	GU831899	GU831899 JN036709 Bacillus thuringiensis strain AIMST KBT9-X		
506	2.2.C	+	GU646907	EU999992 Enterobacter ludwigii strain T4384	Enterobacter	
507		-	GU808439	EU430751 Enterobacter sp. ZJUPD5	Enterobacter	
508	+		GU646889	HQ185399 Stenotrophomonas maltophilia strain 2681	Stenotrophomonas	
511		+	,		Stenotrophomonas	
512		+	GU646901	HQ185399 Stenotrophomonas maltophilia strain 2681	Stenotrophomonas	
523		-	GU808438	EU430751 Enterobacter sp. ZJUPD5	Enterobacter	
524		-	GU831901	HM771092 Bacillus sp. INBio3686F	Bacillus	
539		-	GU831898	JN006263 <i>Bacillus</i> sp. C-21	Bacillus	
546		- GU831892		JN036709 Bacillus thuringiensis strain AIMST KBT9-X	Bacillus	
554		-	GU808442	DQ298127 Paenibacillus polymyxa isolate U4D	Paenibacillus	
555		-	GU808437	HM598440 Enterobacter sp. UFLA81	Enterobacter	
559		-	GU814030	HM355676 Microbacterium foliorum strain BAC3087	Microbacterium	
589		-	GU831897	JN006263 <i>Bacillus</i> sp. C-21	Bacillus	
597		-	GU808434	FN401343 Enterobacter cloacae isolate PHLTA- 11	Enterobacter	
603	4.1.A	-	GU646913	DQ530139 Microbacterium sp. RI48	Microbacterium	

(Table 1) contd....

Isolate	Sampling	Plasmid presence	16S rDNA accession number	First Blast Hit	Phylogenetic affiliation	
613		-	GU814022	HQ877451 Stenotrophomonas sp. 33T	Stenotrophomonas	
616		-	GU808444	AM983496 Paenibacillus sp. AM27T2	Paenibacillus	
619		-	GU808433	JN036709 Bacillus thuringiensis strain AIMST KBT9-X	Bacillus	
620		-	GU831904	FR682931 Stenotrophomonas sp. R-41388	Stenotrophomonas	
625		+	GU646890	EU880530 Paenibacillus sp. PRE17	Paenibacillus	
630		-	GU814033	DQ530139 Microbacterium sp. RI48	Microbacterium	
633		+	GU646891	EU880530 Paenibacillus sp. PRE17	Paenibacillus	
635		+	GU646896	EU880530 Paenibacillus sp. PRE17	Paenibacillus	
636		+	GU646909	EU880530 Paenibacillus sp. PRE17	Paenibacillus	
637		+	GU646892	EU880530 Paenibacillus sp. PRE17	Paenibacillus	
640		+	GU646908	EU880530 Paenibacillus sp. PRE17	Paenibacillus	
659		-	GU646911	FR823407 Bacillus sp. ITCr36	Bacillus	
675		-	GU808430	JN036709 Bacillus thuringiensis strain AIMST KBT9-X	Bacillus	
706		+	GU646893	HM629374 Staphylococcus sp. B-G-R2A2	Staphylococcus	
708		+ GU646894 JF766691 Staphylococcus sp. BIHB 1375		Staphylococcus		
711		-	GU831887	DQ232617 Agromyces sp. VKM Ac-1802	Agromyces	
714		-	GU808441	GU808441 AY337581 Paenibacillus sp. CC-SB818D1		
737		-	GU831911 FM992644 Bacillus safensis strain F5-77		Bacillus	
773		-	GU831886	GU831886 DQ440827 <i>Bosea</i> sp. CRIB-12		
776		- GU831888 FN563149 Rhodococcus equi 103S		Rhodococcus		
777	-		GU831912	JN036709 Bacillus thuringiensis strain AIMST KBT9-X	Bacillus	
796		-	GU808445	JF798384 Paenibacillus lautus strain T1-11	Paenibacillus	
828		-	GU808443	AJ746160 Paenibacillus sp. MG103	Paenibacillus	
829		-	GU831894	JN006263 Bacillus sp. C-21	Bacillus	
836		-	GU831903	DQ026647 Streptomyces exfoliatus strain NRRL B-2494	Streptomyces	
856		-	GU808429	FR667174 Paenibacillus sp. ITP26	Paenibacillus	
860		-	GU646912	AM990746 Bacillus sp. MOLA 522	Bacillus	
874		-	GU831900	FR823407 Bacillus sp. ITCr36	Bacillus	
879	4.1.B	+	GU646902	EU362611 Paenibacillus polymyxa isolate TN99	Paenibacillus	
890		-	GU831902	JF772519 Sinorhizobium sp. bB42(2011)	Sinorhizobium	
901 bis		-	GU831889	GU097198 Paenibacillus sp. AT5	Paenibacillus	
904		-	GU646910	JF798384 Paenibacillus lautus strain T1-11	Paenibacillus	
917		-	GU831893	JN006263 <i>Bacillus</i> sp. C-21	Bacillus	
924		-	GU814031	DQ530139 Microbacterium sp. RI48	Microbacterium	

(Table 1) contd....

Isolate	Sampling	Plasmid presence	16S rDNA accession number	First Blast Hit	Phylogenetic affiliation	
928		-	GU814029	HQ132733 Acinetobacter sp. Ld5	Acinetobacter	
953		-	GU808440	EU430751 Enterobacter sp. ZJUPD5 Entero		
987		- GU831913		JN036709 Bacillus thuringiensis strain AIMST KBT9-X	Bacillus	
1003		-	GU831914	GU831914 GU252111 Bacillus sp. 2008724139		
1005		-	GU831915	JF820106 Bacillus sp. PG-3-9	Bacillus	
1126	4.1.C	-	GU646914	EU729736 Arthrobacter aurescens strain MM10	Arthrobacter	
1128		-	GU646915	HQ597008 Arthrobacter aurescens strain ABRI- INW 23	Arthrobacter	

Symbols: +: presence of plasmid molecules; -: absence of plasmid molecules.

growth rate. Bacteria producing visible colonies at days 1 and 2 (classes 1 and 2) were defined as "fast growers" (copiotrophs or r-strategists), while bacteria that produced colonies later (class 3) were defined as "slow growers" (oligotrophs or K-strategists). The number of bacteria in each class was expressed as a percentage of the total count and gave insight into the distribution of r- and K-strategists in each sample. Characteristics of r-strategists include fast growth in response to medium enrichment, while Kstrategists are characterized by slow growth in response to enrichment.

After six days of growth at 28°C, a set of 1,200 colonies (200 per each of the six soil samples) were randomly chosen from 0.1 TSA plates and isolated on the same medium for the further characterization.

Eco-physiological Index

To express the distribution of the fast- versus slowgrowing bacteria (r- versus K- strategists) in soil samples. the Eco-Physiological (EP) index [12], was calculated using three classes (i.e. colonies grown after 1, 2, and 6 days) [11]. The EP index of each soil tested was calculated using the equation: H'= $-\sum (P_i \times \log_{10} P_i)$, where P_i represents the CFU at each day (1, 2 and 6 days of incubation) as a proportion of the total CFU in that sample after 6 days incubation i.e. the proportion of colonies appearing on counting day i (i = 1, 2, ...6) with $EP_{min} = 0$. Higher values of EP index imply a more even distribution of proportions of bacteria developing on different days (i.e., different classes of bacteria).

Statistics

Bacterial population data (CFU g⁻¹ of soil) were log transformed and subsequently analysed by using t-test (Graph Pad Prism version 5 software). Percentage data of r/K strategists and EP index value were logit-transformed, Logit $(p) = \log [p/(1-p)]$ for the proportion p, and compared using *t*-test (Graph Pad Prism version 5 software).

Agarose Gel Electrophoresis

Agarose gel electrophoresis in TAE buffer (0.04 M Tris-Acetate, 0.01 M EDTA) containing 0.5 µg/ml (w/v) of ethidium bromide [13] was used to check the presence of plasmids (0.8% w/v), and to analyze amplicons obtained either from PCR amplification of 16S rRNA genes (0.8% w/v) or RAPD fingerprinting (2.0 % w/v).

Analysis of Plasmids Content

Analytical amounts of plasmid DNA were obtained from 1.5 ml bacterial cultures using the commercial Kit Plasmid Miniprep (Qiagen) set up for Gram-negative bacteria with the use of a robotic workstation (OiaCube, Oiagen).

PCR Amplification and Sequencing of 16S rRNA Genes

PCR amplification of 16S rRNA genes was carried out according to Papaleo et al. [14] using a MJ Research PTC 100 Peltier Thermal Cycler (CELBIO). Amplicons were excised from agarose gel and purified using the "QIAquick" gel extraction kit (QiAgen). Direct sequencing was performed on both DNA strands using the chemical dye terminator [15].

RAPD Analysis

Random amplification of DNA fragments was carried out using primer 1253 (5' GTTTCCGCCC 3') and the amplification conditions described elsewhere [16].

Homologs Retrieval and Phylogenetic Analysis

Probing of the DNA databases was performed with the BLAST program [17], using default parameters. The ClustalW program [18] was used to align the 16S rRNA gene sequences obtained with the most similar ones retrieved from databases. Each alignment was analyzed using the neighbor-joining method [19] according to the model of Kimura 2-parameter distances [20]. Phylogenetic trees were constructed using the MEGA4 software [21]. The robustness of the inferred trees was evaluated by 1000 bootstrap resamplings.

RESULTS AND DISCUSSION

Characterization of Bacterial Communities Isolated from Soil Samples

Six soil samples were collected from two boxes of UF-NT soil, i.e. from the top (2.2 A, 2.2 B, and 2.2 C samples) and from the bottom (4.1 A, 4.1 B, and 4.1 C samples) of the hillside. The total microbial population density ranged from $Log_{10} 6.01 + 0.05$ (bottom) to $Log_{10} 6.61 + 0.17$ (top) cfu g⁻¹ of soil (Table 2), and no significant difference (P>0.05) was observed between the soil samplings collected from the top and the bottom of the experimental field. The structure of the bacterial soil community at each sub-sampling was investigated using the concept of r/K strategy [11] and reported in Table 3. Results indicated that bacterial colonies visible after one day of incubation (r/K class 1) were more abundant in the bottom soil compared to those in the top soil (P<0.05), whereas no differences (P>0.05) were observed between the bottom and the top samplings colonies visible after two days of incubation (r/K classes 2 and 3) (Table 3). Significant differences in EPI-index between the top and the bottom soil were also found (P<0.05) (Table 3), suggesting changes in community structure of cultivable bacterial communities in the two blocks of experimental field.

Analysis of Plasmid Content

Two hundreds bacterial isolates from each of the six samples (2.2 A, 2.2 B, 2.2 C, 4.1 A, 4.1 B, and 4.1. C) were randomly selected for further characterization. The presence of plasmids was checked as described in Materials and Methods on each of the 1,200 bacterial isolates randomly selected and re-grown on 0.1 TSA medium (Tables 1 and 4). Data obtained are shown in Fig. (2) and revealed that only 27 out of the 1,200 bacterial isolates harbored plasmid molecules. Most of the strains exhibited only a single plasmid molecule, while in a few cases (i.e. isolates 308bis and 363) multiple plasmids were found in the same cell. Furthermore, some isolates showed plasmids with the same electrophoretic

mobility. The size of plasmid molecules ranged between about 2 Kb and 40 Kb, as determined by comparing their eletrophoretic mobility with that of reference plasmids. However, we cannot *a priori* exclude the possibility that the genome of some of the bacterial isolates analyzed might contain large and/or low copy number plasmids, which might have not been revealed by the extraction methodology used in this work.

RAPD Fingerprinting

In order to type the 27 bacterial isolates harboring plasmids, a RAPD [16] analysis using the primer 1253 was carried out. The comparative analysis of RAPD profiles obtained allowed the bacterial strains to be clustered in groups embedding bacterial isolates exhibiting the very same amplification profile (hereinafter haplotype). Bacterial isolates with the same haplotype were considered as the same strain. Data obtained are reported in Fig. (2), which shows that the 27 isolates can be split into 15 RAPD groups. Indeed, some isolates exhibited the same RAPD profile, suggesting that they might correspond to the same bacterial strain. In most cases, isolates exhibiting the same RAPD profile share a plasmid with the same electrophoretic mobility (i.e. the same plasmid if we assume that plasmids with the same electrophoretic mobility correspond to the same molecule) (see, for instance, isolates 230, 251, 253, 327 and 398 – RAPD haplotype 3). Just in one case, isolates sharing the same RAPD profile (i.e. cells of the same strain) harbored different plasmids (isolates 457, 500, 506 - RAPD haplotype 13), suggesting that the same strain may host different plasmids.

Table 2. Bacteria colony counts (Log cfu g⁻¹ of soil) as they appeared on 0.1 TSA over a period of six days and total culturable bacteria.

Soil samples	amples r/K class 1 ^a r/K class 2 ^a		r/K class 3ª	Total culturable	
	r-strat	tegists ^b	K-strategists ^b	bacteria	
Тор					
2.2 A	5.90	5.31	5.85	6.23	
2.2 B	6.32	5.98	6.02	6.61	
2.2 C	5.61	5.42	5.59	6.03	
Mean <u>+</u> SEM	5.94 <u>+</u> 0.21 °	5.57 <u>+</u> 0.21 °	5.82 <u>+</u> 0.12 °	6.29 <u>+</u> 0.17 °	
Bottom					
4.1 A	5.91	5.14	5.59	6.13	
4.1 B	5.80	5.20	5.36	6.01	
4.1 C	5.80	5.03	5.24	5.96	
Mean <u>+</u> SEM	5.84 <u>+</u> 0.04 °	5.12 <u>+</u> 0.05 °	5.40 <u>+</u> 0.10 °	6.03 <u>+</u> 0.05 °	
Total Mean <u>+</u> SEM	5.89 <u>+</u> 0.10	5.35 <u>+</u> 0.14	5.61 <u>+</u> 0.12	6.16 <u>+</u> 0.10	

a r/K class 1 = bacteria producing visible colonies at day 1

r/K class 2 = bacteria producing visible colonies at day 2

r/K class 3 = bacteria producing visible colonies at day 6

Bacteria recovered in day 1 and 2 are fast growers (r-like strategists) and those recovered on day 6 are slow growers (K-like strategists).

Log data were analyzed for statistical significance by using t-test (Graph Pad Prism version 5 software). Values \pm SEM followed by the same letter are not significantly different (P > 0.05) within each vertical column.

Table 3. Percentages of r/K bacterial strategists (classes 1, 2 and 3) on total culturable bacteria colony counts as they appeared on 0.1 TSA over a period of six days and EPI-index.

Soil samples	Soil samples r/K class 1 r/K class 2		r/K class 3	EPI-index	
	r-stra	tegists ^a	K-strategists ^a		
Тор					
2.2 A	46%	12%	42%	0.42	
2.2 B	51%	23%	26%	0.45 0.47	
2.2 C	39%	25%	37%		
Mean <u>+</u> SEM	45.33 <u>+</u> 3.48 ^b	20.00 <u>+</u> 4.04 ^a	35.00 <u>+</u> 4.73 ^a	0.47 <u>+</u> 0.01 ^b	
Bottom					
4.1 A	61%	10%	29%	0.39	
4.1 B	62%	15%	23%	0.40	
4.1 C	69%	12%	19%	0.36	
Mean + SEM	64.00 ± 2.52 ^b	12.33 <u>+</u> 1.45 ^a	23.67 ± 2.91 ^a	0.41 ± 0.02 ^b	

^a Bacteria recovered in day 1 and 2 are fast growers (r-like strategists) and those recovered on day 6 are slow growers (K-like strategists).

Phylogenetic Affiliation of Bacterial Isolates Harboring **Plasmid Molecules**

To affiliate each bacterial isolate to a given taxon, the nucleotide sequence of the 16S rRNA gene from at least one isolate per each RAPD group was determined. The 16S rRNA genes were PCR-amplified and sequenced from 25 isolates and their analysis revealed that:

- i) The 15 RAPD haplotypes were representative of seven bacterial genera, two Gram positive (Staphylococcus and Paenibacillus) and five Gram negative (Acinetobacter, Enterobacter, Pantoea, Stenotrophomonas, and Klebsiella, all belonging to γ-proteobacteria), with Acinetobacter and Paenibacillus representing half of the bacteria-harboring plasmids community (Fig. 2 and Table 4).
- ii) The 16S rRNA gene sequences from three Enterobacter isolates (457, 500, 506) were identical in agreement with the finding that they also share the same RAPD profile. The fourth Enterobacter isolate (363), exhibiting a different RAPD profile (Fig. 2), also possesses a 16S rRNA gene sequence differing in one position in respect to the other three ones.
- iii) The two Staphylococcus strains (706 and 708 exhibiting different RAPD haplotypes) shared the same 16S rRNA gene sequence, suggesting that they belong to the same species.
- iv) The Acinetobacter sequences were placed in three distant branches of the trees, suggesting that they very likely belong to (at least) three different species, a finding that is in agreement with their very different RAPD profile.

Analysis of the Composition of the Aerobic Heterotrophic Cultivable Bacterial Communities Lacking Small **Plasmids**

In order to get some information also on the composition of the heterotrophic cultivable bacterial community, which did not exhibit plasmids under the experimental conditions used in this work and to compare it with the taxonomical position of bacteria harboring small plasmids, the 16S rRNA genes were amplified and sequenced as described in Materials and Methods from a panel of 79 randomly chosen bacterial isolates (Table 1). The analysis of the 79 nucleotide sequences obtained revealed that they were affiliated to 14 bacterial genera, with Bacillus (21 isolates) and Stenotrophomonas (15 isolates) being the most represented ones. Half of the isolates belong to Gram- bacteria represented only by members of the α - and γ -proteobacteria; the other isolates belong mainly to Bacillus and Paenibacillus, even though representatives of different genera (Agromyces, Arthrobacter, Microbacterium, Rhodococcus, and Streptomyces) of high GC Gram+ bacteria were disclosed.

Data reported in Table 4 revealed that bacteria belonging to Stenotrophomonas, Acinetobacter, Enterobacter and Paenibacillus included both isolates harboring or lacking plasmid molecules. To check the existence of a possible correlation between the presence of plasmids and the phylogenetic position of bacterial isolates, a phylogenetic tree for each of these four genera was constructed (Fig. 3), whose analysis revealed that some isolates embedded in the same genus very likely belong to different species, since the sequences joined different clusters of a phylogenetic tree. This is particularly true for Enterobacter and Acinetobacter isolates, whereas Stenotrophomonas and Paenibacillus exhibited a more homogeneous distribution within the respective tree.

b Percentage data were Logit transformed and analyzed for statistical significance by using t-test (Graph Pad Prism version 5 software). Values ± SEM followed by the same letter are not significantly different (P > 0.05) within each vertical column. Significant data are also indicated in bold.

Table 4. Number of bacterial isolates belonging to different taxonomic groups recovered from the different soil samplings.

			Soil sample							
		Plasmid	Тор			Bottom		_		
Organism	Taxonomy		2.2.A	2.2.B	2.2.C	4.1.A	4.1.B	4.1.C	Sub total	Total
				I	Isolate	e numbering	I.			
			1-200	201-400	401-600	601-800	801-1000	1001-1200		
Stenotrophomonas		+	0	0	4	0	0	0	4	- 19
Sienotrophomonas		-	5	7	1	2	0	0	15	17
Acinetobacter		+	0	7	1	0	0	0	8	13
Acmeiobacier		-	0	3	1	0	1	0	5	13
Enterobacter	γ - proteobacteria	+	0	1	3	0	0	0	4	13
Enterobacter		-	0	2	6	0	1	0	9	13
Pseudomonas		+	0	1	0	0	0	0	1	1
Pantoea		+	0	0	1	0	0	0	1	1
Klebsiella		-	1	0	0	0	0	0	1	1
Agrobacterium		-	0	1	0	0	0	0	1	1
Bosea	α - proteobacteria	-	0	0	0	1	0	0	1	1
Sinorhizobium		-	0	0	0	0	1	0	1	1
Paenibacillus	0 1 0.0	+	0	0	0	6	1	0	7	16
Faenibaciiius	Gram+ low G+C	-	1	0	1	3	4	0	9	
Staphylococcus		+	0	0	0	2	0	0	2	2
Bacillus	Gram + low GC	-	3	0	6	5	5	2	21	21
Agromyces		-	0	0	0	1	0	0	1	1
Streptomyces		-	1	0	0	0	1	0	2	2
Rhodococcus	Gram+ High GC	-	1	0	0	1	0	0	2	2
Arthrobacter		-	0	0	0	0	0	2	2	2
Microbacterium		-	2	1	3	2	1	0	9	9
	Total Isolates +			9	9	8	1	0		27
				14	18	15	14	4		79
Total				18			9			27
	-		46			33			79	

Symbols: +/- represent the presence/absence of plasmids, respectively.

CONCLUSION

In this work we have analyzed the aerobic soil heterotrophic cultivable bacterial community consisting of 1,200 bacterial isolates from NT and UF soil, with high organic input and increased enzyme activities, which favour functional diversity of the microbial community [10]. Total plate counts did not reveal any differences between the two subsamplings (i.e. top and the bottom of the experimental field), while significant differences were found in the structure of

the soil bacterial community that showed an uneven distribution of r- and K- strategists in the two sub-samples. Concerning the presence and the frequency of plasmids in these communities, data obtained suggest that they were harbored only by a low percentage (2.0 %) of bacterial isolates, that their size ranged between 2 and 40 kb, and that multiple plasmids were present only in a very limited number of isolates. The lack of plasmids of higher size did not *per se* imply their absence in the bacterial cells analyzed in this work;

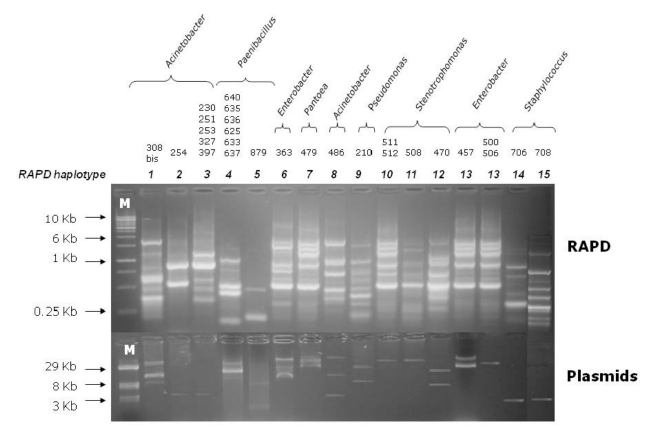


Fig. (2). Agarose gel electrophoresis of RAPD profiles (upper) from 27 soil bacterial isolates harboring plasmids (lower). The last line of the right side was cut from another figure and pasted in Fig. (2), without changing neither the intensity nor the size of each band. Line M: (upper) DNA linear marker; (lower) reference plasmids of known size.

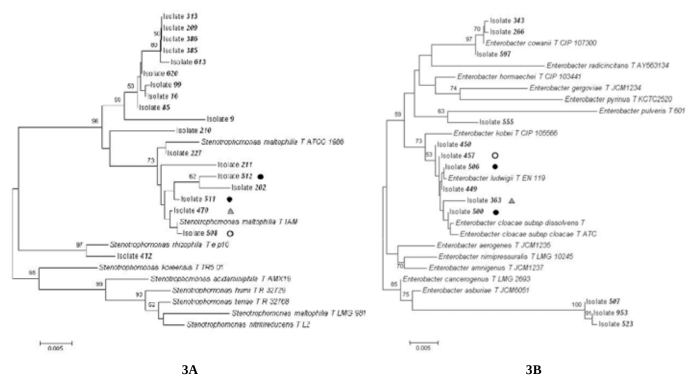


Fig. (3) contd....



Fig. (3). Phylogenetic trees constructed using the 16S rRNA sequences obtained from *Stenotrophomonas* (**A**), *Enterobacter* (**B**), *Paenibacillus* (**C**), and *Acinetobacter* (**D**) isolates analyzed in this work and the most similar sequences retrieved from databases. Bootstrap values > 50 are shown. The pairwise deletion option was used. Isolates harboring one or more plasmids are marked by a black or white dot or a grey triangle. Isolates harboring a plasmid with the same electrophoretic mobility are marked with the same symbol.

indeed, the procedure for plasmid extraction utilized in this work did not allow the isolation of plasmid molecules with a size higher than 40 kb.

We are completely aware that the size of the bacterial communities (1,200 isolates) that we have analyzed is much smaller of the extant total soil microbial communities and thus it represent a small sub-fraction of it. In spite of this limitation, the overall phylogenetic analysis revealed an uneven distribution of both species and plasmid-harboring strains. Interestingly, in most cases, isolates harboring plasmids of the same or different length clustered together in the same branch of the tree, which however also includes strains lacking plasmids. This might suggest that plasmids preferentially flow (vertically and/or horizontally) between (closely) related strains rather than between bacteria belonging to different species of the same genus. This is the case of Enterobacter, where at least three different strains of the same species show the same plasmid profile. Data reported in Table 4 also revealed that there was not a uniform distribution of plasmid-harboring isolates in the two samples collected from the top and the bottom of the experimental field. Indeed, Stenotrophomonas, Acinetobacter and Enterobacter isolates were detected only in the first sub-sample, whereas the Paenibacillus isolates were disclosed in the second one. Furthermore, some bacterial genera (such as Bacillus spp.) did not exhibit any plasmid molecule, even though they were "over"-represented in the bacterial community.

CONFLICT OF INTEREST

The authors confirm that this article content has no conflict of interest.

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