High taxonomic level fingerprint of the human intestinal microbiota by Ligase Detection Reaction - Universal Array approach

Marco Candela*1, Clarissa Consolandi2, Marco Severgnini2, Elena Biagi1, Bianca Castiglioni3, Beatrice Vitali1, Gianluca De Bellis2 and Patrizia Brigidi1

Background: Human beings have been recently reconsidered as super-organisms in co-evolution with an immense microbial community living in the gastrointestinal tract (GIT), the human intestinal microbiota [1,2]. Providing important metabolic functions that we have not evolved by our own [3], the intestinal microbiota has a fundamental role for the human health and well being [4,5]. Several of our physiological features, such as nutrient processing, maturation of the immune system, pathogen resistance, and development of the intestinal architecture, strictly depend on the mutualistic symbiotic relationship with the intestinal microbiota [6]. On the basis of its global impact on human physiology, the intestinal microbiota has been considered an essential organ of the human body [7].

The composition of the adult intestinal microbiota has been determined in three large scale 16S rRNA sequences surveys [7-11]. The phylogenetic analysis of a total of 45,000 bacterial 16S rRNA data from 139 adults revealed that, at the phylum level, only a small fraction of the known bacterial diversity is represented in our GIT. The vast majority of bacteria in the human intestinal microbiota (>99%) belongs to six bacterial phyla: Firmicutes, Bacteroidetes, Actinobacteria, Proteobacteria, Verrucomicrobia and Fusobacteria. The two dominant divisions are Firmicutes and Bacteroidetes, which represent together up to 90% of the total microbiota, with a relative abundance of 65% and 25%, respectively. Actinobacteria, Proteobacteria, Verrucomicrobia and Fusobacteria are the subdominants phyla with a relative abundance up to 5, 8,
2 and 1%, respectively. On the contrary, at lower taxonomic levels, we assist to a real explosion of the bacterial diversity in the human GIT. At least 1,800 genera [≥ 90% of sequence identity (ID)] and 16,000 phylotypes at the species level (≥ 97% ID) have been identified until now, predicting even a greater diversity at the species level [8]. Since 70% of these phylotypes are subject-specific, and no phylotype is present at more than 0.5% abundance in all subjects [12], the intestinal microbiota of each individual has been shown to consist in a subject specific complement of hundreds of genera and thousands of species. However, the large degree of functional redundancy between species and genera allowed identifying a core microbiome at the gene level which is shared between all individuals [12]. Coding for genes involved in important metabolic functions, this core functional microbiome is fundamental to support the mutualistic symbiotic relationship with the human host.

Recently, 16S rRNA sequences studies have been carried out with the attempt to describe disease-associated imbalances of the human intestinal microbiota. Even though species variability was associated with inter-individual variability, phylum-level changes of the intestinal microbiota were associated with specific diseases. In particular, obesity was characterized by a higher proportion of Firmicutes and Actinobacteria with respect to Bacteroidetes and an overall reduced bacterial diversity [12,13]. Differently, inflammatory bowel diseases (IBD) were characterized by a marked reduction of bacterial diversity in the Clostridium cluster IV and XIVa belonging to Firmicutes, a decline in Bacteroidetes biodiversity, and a correspondent increase in Proteobacteria and Bacillus [14,15]. Analogously, intestinal inflammation has been generally related with a marked increase in Enterobacteriaceae and a correspondent decrease in members of the resident colonic bacteria [16,17]. In the light of these findings, it has been recently hypothesized that these high level taxonomic imbalances of the human intestinal microbiota can cause deviations from the core functional microbiome with a final impact on the host physiological state [12,18,19].

Since more than 75% of the phylotypes detected in the human GIT does not correspond to cultured species [20], phylogenetic DNA-microarrays have been recognized as a valuable tool for a high-throughput, quantitative and systematic analysis of the human intestinal microbiota [21]. Recently, three different small ribosomal subunit RNA (SSU rRNA) based high-density phylogenetic microarrays for studying the human microbiota have been developed [22-24]. Targeting thousands bacterial phyo-types, these DNA-microarrays have been successfully applied in studies for the deep phylogenetic characterization of the human intestinal microbiota.

In order to specifically monitor the microbiota imbalances that impact on human physiology independently of the inter-individual variability, here we developed an original DNA-microarray for the high taxonomic level fingerprint of the human intestinal microbiota, called HTF-Microbi.Array (High Taxonomic Fingerprint Microbiota Array). The relatively low number of targets allowed implementing the Ligase Detection Reaction (LDR) technology [25,26] for the development of the HTF-Microbi.Array. This enzymatic in vitro reaction, based on the discriminative properties of the DNA ligase enzyme, requires the design of a pair of two adjacent oligonucleotides specific for each target sequence: a probe specific for the variation (called "Discriminating Probe", or DS) which carries a 5’-fluorescent label, and a second probe, named "Common Probe" (or CP), starting one base 3’-downstream of the DS that carries a 5’-phosphate group and a unique sequence named cZipCode at its 3’-end. The oligonucleotide probe pairs and a thermostable DNA ligase are used in a LDR reaction with previously PCR-amplified DNA fragments. This reaction is cycled to increase product yield. The LDR products, obtained only in presence of a perfectly matching template by action of the DNA ligase, are addressed to a precise location onto a Universal Array (UA), where a set of artificial sequences, called Zip-codes are arranged. These products carry both the fluorescent label and a unique cZipCode sequence and can be detected by laser scanning and identified according to their location within the array. The LDR approach is a highly specific and sensitive assay for detecting single nucleotide variations; thus, differences of a single base along the 16S rRNA gene can be employed to distinguish among different microbial lineages. The HTF-Microbi.Array was successfully tested in a pilot study for the characterization of the faecal microbiota of eight healthy young adults.

**Results**

**Target selection and probe design**

The rational selection of the HTF-Microbi.Array targets was carried out using a phylogenetic approach. To this aim we implemented the 16S rRNA database of the ARB Project (release February, 2005) with the 16S rRNA gene database of the RDP available at the time and a phylogenetic tree was constructed. Based on the tree nodes, 30 phylogenetical groups of the human intestinal microbiota were rationally selected as the target group for the HTF-Microbi.Array. In Fig. 1 we report the phylogenetic tree of the 16S rRNA sequences of the HTF-Microbi.Array positive set. The selected groups belonged to different phylogenetic levels (species, genus, family, cluster, or group of species indicated by the wording "et rel."). The entire list of the array targets is represented in Table 1. For part of the division Firmicutes, the target
Figure 1 SSU rRNA based phylogenetic tree of the 16S rRNA sequences of the HTF-Microbi.Array positive set. For each node we report the number of sequences used from our ARB 16S rRNA sequence database. The triangles dimension is proportional to the number of sequences clustered together. The phylogenetic tree was obtained by using the neighbour-joining algorithm for the sequence alignment in ARB software.
### Table 1: Probe set of the HTF-Microbi.Array.

| PROBE                  | N. | TAXONOMIC LEVEL | CLUSTER | ORDER | DIVISION | ECO | H.G. AB % |
|------------------------|----|-----------------|---------|-------|----------|-----|-----------|
| Bacteroides/Prevotella | 16 | Cluster         | Bacteroides/Prevotella | Bacteroides | Bacteroidetes | M   | 20        |
| Ruminococcus bromii    | 38 | Sub cluster     | Cl IV   | Clostridiales | Firmicutes | M   |           |
| Ruminococcus albus     | 39 | Sub cluster     | Cl IV   | Clostridiales | Firmicutes | M   |           |
| Faecalibacterium praunstizii | 40 | Sub cluster     | Cl IV   | Clostridiales | Firmicutes | M   |           |
| Oscillospira guillermondii | 41 | Sub cluster     | Cl IV   | Clostridiales | Firmicutes | M   | 65        |
| Clostridium IX         | 37 | Cluster         | Cl IX   | Clostridiales | Firmicutes | M   |           |
| Veillonella            | 20 | Species (et rel)| Cl IX   | Clostridiales | Firmicutes | M   |           |
| Clostridium XIVa       | 22 | Cluster         | Cl XIVa | Clostridiales | Firmicutes | M   |           |
| Eubacterium rectale    | 19 | Species (et rel)| Cl XIVa | Clostridiales | Firmicutes | M   |           |
| Bifidobacteriaceae     | 25B| Family          | Bifidobacterium | Bifidobacteriales | Actinobacteria | M   | 5         |
| B. longum              | 3  | Species (et rel)| Bifidobacterium | Bifidobacteriales | Actinobacteria | M   |           |
| Lactobacillaceae       | 21B| Family          | Lactobacillaceae | Lactobacillales | Firmicutes | M   |           |
| L. plantarum           | 33 | Species (et rel)| Lactobacillaceae | Lactobacillales | Firmicutes | M   | <1        |
| L. casei               | 12 | Species (et rel)| Lactobacillaceae | Lactobacillales | Firmicutes | M   |           |
| L. salivarius          | 14 | Species (et rel)| Lactobacillaceae | Lactobacillales | Firmicutes | M   |           |
| Bacillus clausii       | 32 | Species (et rel)| Bacillaceae | Bacillales | Firmicutes | M   | <1        |
| Bacillus subtilis      | 8  | Species (et rel)| Bacillaceae | Bacillales | Firmicutes | M   | <1        |
| Fusobacterium          | 15 | Genus           | Fusobacteriaceae | Fusobacteria | Fusobacteria | M   | <0.5      |
| Cyanobacteria          | 42 | Family          | Cyanobacteria | Cyanobacteria | Cyanobacteria | M   | <0.1      |
| Clostridium XI         | 36 | Cluster         | Cl XI   | Clostridiales | Firmicutes | O   | 0         |
| Clostridium difficile  | 18 | Species (et rel)| Cl XI   | Clostridiales | Firmicutes | O   |           |
| Clostridium I and II   | 35 | Cluster         | Cl I and II | Clostridiales | Firmicutes | O   | 0         |
| Clostridum perfringens | 17 | Species (et rel)| Cl I and II | Clostridiales | Firmicutes | O   |           |
| Enterococcus faecalis  | 9  | Species (et rel)| Enterococcales | Lactobacillales | Firmicutes | O   | <1        |
| Enterococcus faecium   | 10 | Species (et rel)| Enterococcales | Lactobacillales | Firmicutes | O   | <1        |
selection was carried out based on the classification proposed by Collins et al. [27]. *Clostridium* cluster I and II, *Clostridium* cluster IX, *Clostridium* cluster XI, and *Clostridium* cluster XIVa were selected. For the *Clostridium* cluster IV, four subgroups of species were defined: *Ruminococcus albus et rel*, *Ruminococcus bromii et rel*, *Faecalibacterium prausnitzii et rel*., and *Oscillospira guillermondii et rel*. Within the *Firmicutes* division, the family *Lactobacilliaceae*, and the groups *Bacillus clausii et rel.*, *Bacillus subtilis et rel.*, *Bacillus cereus et rel.*, *Enterococcus faecalis et rel.*, and *Enterococcus faecium et rel.* were also selected. Other selected groups were the *Bacteroides/Prevotella* cluster (division *Bacteroides*), the family *Bifidobacteriaceae* (division *Actinobacteria*), the family *Enterobacteriaceae* and the genus *Campylobacter* (division *Proteobacteria*). For clusters or families, relevant species, genera or subgroups of species were selected to design "sub-probes". The genus *Veillonella* was selected for *Clostridium* cluster IX, the species *Eubacterium rectale* for *Clostridium* cluster XIVa, *Clostridium difficile* for *Clostridium* cluster XI, and *Clostridium perfringens* for *Clostridium* cluster I and II. The group *Bifidobacterium longum et rel.* was chosen for the family *Bifidobacteriaceae*, and the genera *Yersinia* and *Proteus* for the *Enterobacteriaceae*. Based on an original phylogenetic design, the entire probe set of the HTF-Microbi.Array cover up to 95% of the bacterial groups belonging to the human intestinal microbiota [28].

Specificity and coverage of each candidate probe was assessed by using the tool Probe Match of the RDP database. The probe pairs selected for the HTF-Microbi.Array were required to perfectly match the sequences of the positive set and to possess at least a mismatch at the 3' end of the discriminating probe respect to the entire negative set. The designed probes pairs had an average melting temperature (Tm) of 67.8 ± 0.9°C (n = 60) and an average length of 35.6 ± 4.9 nucleotides. Sixteen out of the 30 probe pairs were characterized by having no degenerated bases, whereas only one probe pair (i.e. the one for *Clostridium* cluster I and II) had 4 and 3 ambiguous bases on DS and CP, respectively (Additional file 2).

**Validation of the HTF-Microbi.Array**

**LDR probe pair specificity**

The specificity of the designed LDR probe pairs was tested by using 16S rRNA PCR amplicons from 28 microorganisms members of the human intestinal microbiota. Amplicons were prepared by amplification of genomic DNA extracted from DSMZ cultures or genomic DNA from ATCC collection. Proving the specificity of the HTF-Microbi.Array all the 16S rRNA amplicons were properly recognized in separate LDR hybridization reactions with the entire probe set of the array. Two replicated independent LDR-UA experiments were performed with an optimal reproducibility (Additional file 3). For each of the 16S rRNA template only group-specific spots, and spots corresponding to the hybridization controls showed positive signals (*P < 0.01*) (Table 2). As a negative control, we performed two independent PCR-LDR-UA experiments using double distilled water, instead of genomic DNA, as sample. As expected, no positive signal was detected. The ratio between the signal intensities of the specific probes and the blank intensity (SNR,) averaged 206.9 ± 185.7, whereas the ratio between all the other probes and the blank intensity (SNR,blank) averaged 2.1 ± 1.4. Therefore, the ratio between specific and non-specific probes resulted more than 100 fold on average.

**Evaluation of the LDR sensitivity and relative abundance detection level**

In order to define the detection limits of the HTF-Microbi.Array, LDR-UA experiments were carried out with different concentrations of an artificial mix of 16S rRNA amplicons from 6 members of the human intestinal microbiota. The 16S rRNA amplicons from *Bacillus cereus*, *Lactobacillus casei*, *Bifidobacterium adolescentis*, *Ruminococcus albus*, *Prevotella*, *Y. enterocolitica* were all specifically recognized in a range of concentrations from 0.7 to 75 fmol (*P < 0.01*), demonstrating the high sensitivity and specificity of the array (Fig. 2). Subsequently, in

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**Table 1: Probe set of the HTF-Microbi.Array. (Continued)**

| Genus/Species | Species (et rel) | Family | Tribe | Order | Class | Phyla | Probe | P | O/P | SNR | SNRblank |
|---------------|------------------|--------|-------|-------|-------|-------|-------|-----|-----|-----|-----------|
| Enterobacteriaceae | 238 | Family | Enterobacteriaceae | Enterobacterales | Proteobacteria | O/P | <8 | 0 |
| Yersinia enterocolitica | 4 | Species (et rel) | Enterobacteriaceae | Enterobacterales | Proteobacteria | O/P | 0 |
| Proteus | 5 | Genus | Enterobacteriaceae | Enterobacterales | Proteobacteria | O/P | 0 |
| Campylobacter | 6 | Genus | Campylobacteraceae | Campylobacterales | Proteobacteria | P | 0 |

For each probe is indicated the spot number, the phylogenetic level, the phylogeny of the target group, the ecology in the gastrointestinal ecosystem (mutualistic (M), opportunistic (O), pathogen (P)). The relative abundance in a healthy gut ecosystem of the principal microbial groups is also indicated.
### Table 2: Specificity test.

| DNA Target            | Positive signal                  | SNR other | SNR spec | p-value spec |
|-----------------------|----------------------------------|-----------|----------|--------------|
| *B. fragilis* ATCC25285 | *Bacterodes/Prevotella*          | 0.85      | 30.81    | 9.35E-05     |
|                       |                                  | 0.53      | 21.45    | 7.39E-04     |
| *B. thetaiotaomicrom* ATCC29143 | *Bacterodes/Prevotella*          | 0.45      | 61.44    | 2.56E-04     |
|                       |                                  | 1.66      | 347.24   | 9.10E-06     |
| *L. gasseri* DSM20243     | *Lactobacillaceae*               | 0.30      | 5.58     | 4.98E-03     |
|                       |                                  | 1.56      | 20.59    | 6.58E-03     |
| *P. melaninogenica* ATCC25845 | *Bacterodes/Prevotella*          | 1.54      | 480.24   | 6.02E-08     |
|                       |                                  | 0.90      | 266.63   | 3.74E-09     |
| *B. subtilis* DSM704      | *Bacillus subtilis*              | 7.93      | 637.39   | 1.56E-09     |
|                       |                                  | 5.62      | 350.10   | 1.47E-05     |
| *E. coli* ATCC11105      | *Enterobacteriaceae*             | 3.27      | 555.04   | 8.65E-08     |
|                       |                                  | 2.59      | 222.39   | 4.50E-07     |
| *P. mirabilis* DSM4479    | *Proteus*                        | 2.42      | 703.22   | 7.74E-09     |
|                       | *Enterobacteriaceae*              | 2.03      | 497.10   | 1.97E-09     |
| *B. bifidum* DSM20456    | *Bifidobacteriaceae*             | 2.67      | 289.39   | 4.78E-11     |
|                       |                                  | 2.23      | 407.10   | 2.40E-08     |
| *L. casei* DSM20011      | *Lactobacillaceae, L. casei*     | 2.59      | 125.13   | 1.01E-04     |
|                       |                                  | 2.26      | 134.78   | 5.92E-04     |
| *Y. enterocolitica* (faecal isolate) | *Yersinia enterocolitica, Enterobacteriaceae* | 1.53      | 231.33   | 1.01E-05     |
|                       |                                  | 2.89      | 340.20   | 1.61E-06     |
### Table 2: Specificity test (Continued)

| Species                  | Genus             | Species Name               | Growth Rate | Concentration | p-value  |
|--------------------------|------------------|-----------------------------|-------------|---------------|----------|
| B. cereus DSM31          | Bacillus cereus  |                             | 2.83        | 193.85        | 1.53E-06 |
|                          |                  |                             | 2.49        | 196.82        | 4.16E-03 |
| B. adolescentis ATCC15703| Bifidobacteriaceae|                            | 4.10        | 732.95        | 3.95E-10 |
|                          |                  |                             | 2.90        | 338.59        | 5.59E-07 |
| L. ramnosus DSM20021     | Lactobacillaceae, L. casei |              | 2.40        | 101.76        | 1.41E-03 |
|                          |                  |                             | 4.23        | 177.70        | 4.62E-07 |
| L. delbrueckii DSM20074  | Lactobacillaceae |                             | 3.77        | 210.11        | 2.24E-08 |
|                          |                  |                             | 3.10        | 121.93        | 6.27E-08 |
| L. pentosus DSM20314     | Lactobacillaceae |                             | 3.05        | 131.65        | 4.58E-09 |
|                          |                  |                             | 1.63        | 58.30         | 5.32E-07 |
| L. acidophilus DSM20079  | Lactobacillaceae |                             | 2.39        | 68.49         | 8.70E-05 |
|                          |                  |                             | 2.66        | 78.50         | 5.88E-06 |
| L. reuteri DSM20016      | Lactobacillaceae |                             | 3.17        | 150.57        | 4.66E-09 |
|                          |                  |                             | 1.74        | 83.60         | 1.98E-07 |
| L. plantarum DSM21074   | Lactobacillaceae, L. plantarum |           | 2.12        | 197.32        | 3.79E-09 |
|                          |                  |                             | 2.09        | 148.35        | 2.77E-08 |
| C. difficile ATCCBAA1382 | Clostridium XI, Clostridium difficile |       | 1.12        | 238.87        | 4.88E-04 |
|                          |                  |                             | 0.80        | 126.38        | 1.96E-03 |
| C. jejuni ATCC33292      | Campylobacter jejuni |                          | 0.70        | 19.89         | 5.29E-03 |
|                          |                  |                             | 0.91        | 28.44         | 5.69E-03 |
order to evaluate the relative abundance detection level of the HTF-Microbi.Array, LDR-UA experiments were performed on hybridization mixes containing low quantities of *Escherichia coli* PCR products and increasing amounts of human genomic DNA. This is a fundamental issue in the case of single species present in the gut microbiota at very low fractional abundance (< 0.1%) [21]. According to our data, 1 fmol of *E. coli* amplicon was sufficient (p < 0.005) to be detected in all the tested conditions (from up to 6.3 μg of human gDNA) (Additional file 4). Considering the PCR product as a ~1700 bp amplicon, 1 fmol corresponds to 1.2 ng and, thus, the sensitivity limit results 0.02%.

| Organism                  | Family, Genus/Species        | SNR  | SNRns       | p-value  |
|---------------------------|-------------------------------|------|-------------|----------|
| *V. parvula* ATCC10790    | Veillonella, Clostridium IX   | 1.12 | 205.66      | 1.57E-04 |
|                           |                               | 0.99 | 140.95      | 1.39E-04 |
| *B. breve* DSM20091       | *Bifidobacteriaceae*          | 2.22 | 570.01      | 6.22E-05 |
|                           |                               | 1.69 | 289.07      | 2.72E-04 |
| *B. longum* ATCC15707     | *Bifidobacteriaceae, B. longum* | 1.76 | 341.94      | 1.64E-03 |
|                           |                               | 0.66 | 134.86      | 4.26E-02 |
| *R. productus* ATCC23340  | *Clostridium XIV*             | 0.64 | 4.21        | 1.41E-03 |
|                           |                               | 1.06 | 17.16       | 1.24E-06 |
| *L. salivarius* SV2       | *Lactobacillaceae, L. salivarius* | 0.89 | 12.23       | 4.34E-04 |
|                           |                               | 0.65 | 7.27        | 2.69E-05 |
| *E. faecalis* ATCC700802  | *E. faecalis*                 | 3.12 | 306.51      | 1.09E-03 |
|                           |                               | 2.27 | 217.16      | 6.56E-03 |
| *C. leptum* DSM73         | *Ruminococcaceae bromii*      | 2.28 | 88.89       | 5.52E-07 |
|                           | *Clostridium IV*              | 1.13 | 39.86       | 2.00E-07 |
| *R. albus* DSM20455       | *Ruminococcus albus*          | 1.46 | 47.05       | 2.50E-07 |

Table reporting the results of the tests to assess probe specificity: 28 bacterial DNA targets were chosen to validate the probe pairs. For each DNA analyzed we report: probe pair showing significant signals, SNR, SNRns (see main text for acronym definitions). The p-values of specific probes are reported for each duplicate experiment. Where needed (i.e. more than one probe pair was present), data are the average of the positive signals (for both SNR, and p-values).
Characterization of the faecal microbiota of eight healthy young adults

The HTF-Microbi.Array was applied in a pilot study for the characterization of the faecal microbiota of eight young adults. For all subjects faecal DNA was extracted, total bacterial 16S rRNA amplified, and two separate LDR-UA experiments were carried out (Additional file 5). For each sample a profile of presence-absence probes response was obtained. The cluster analysis of the phylogenetic fingerprints showed that, with the exception of subject n. 2, samples from the same subject clustered together. The reproducibility of the experiments was evaluated by considering the percentage of the probes giving the same response in both the technical replicates of each sample. With the exclusion of subject n. 2, an average reproducibility of 96% was obtained for all the subject under study, demonstrating a good reproducibility of the microbiota fingerprints obtained using the HTF-Microbi.Array (Fig. 3). As expected, the major mutualistic symbionts of the human intestinal microbiota, such as Bacteroidetes and the members of the Clostridium cluster IV and XIVa, were represented in the faecal microbiota of all the subjects. With the exception of B. clausii et rel., minor mutualistic symbionts such as Actinobacteria, Lactobacillaceae, B. subtilis et rel., Fusobacterium, and Cyanobacteria were detected only in different sub-fractions of the subjects. In particular, subjects n. 17, 15, 4, and 1 were characterized by the presence of Fusobacterium. Subjects n. 4, 15 and 17 possessed B. subtilis et rel., while subjects n. 4, 1, 9, 16 and 5 harboured Cyanobacteria in their faecal microbiota. On the other hand, only a fraction of the subjects, clustering on the left side of the map, presented opportunistic pathogens in their faecal microbiota. Subjects n. 17, 15 and 4 presented both Proteus and E. faecalis et rel., while in subject n. 15 members of the Clostridium cluster I and II and Yersinia...
et rel. were also detected. For each subject the relative fluorescence intensity (IF) contribution of each HTF-Microbi.Array probes, in terms of percentage of the total IF, was also calculated (Fig. 4). The mean of IF data from both the LDR-UA experiments were considered. Even if all subjects were characterized by a specific individual profile, a common trend can be found by comparing the comprehensive relative IF contribution of probes targeting major mutualistic symbionts (Bacteroides/Prevotella, Clostridium clusters IV, IX, and XIVa), minor mutualistic symbionts (Bifidobacteriaceae, Lactobacillaceae, B. clausii et rel., B. subtilis et rel., Fusobacterium, and Cyanobacteria), and opportunistic pathogens (Clostridium clusters I and II, X, E. faecalis et rel., E. faecium et rel., B. cereus et rel., Enterobacteriaceae, Yersinia, Proteus, Campylobacter). In particular, for all subjects the highest relative IF contributions were obtained for major mutualistic symbionts. The contribution of Bacteroides/Prevotella ranged between 8-37%, whereas the contribution of Clostridium clusters IV, IX, and XIVa ranged between 17-34%, 3-15%, and 5-29%, respectively. Differently, minor mutualistic symbionts were characterized by lower values of relative IF contributions. Bifidobacteriaceae contributed for the 0.5-3.1%, Lactobacillaceae for the 1.5-9.4%, B. clausii et rel. for the 4-13%, B. subtilis et rel. for the 0.6-2.5%, Fusobacterium for the 1.2-4.4%, and Cyanobacteria for 0.6-4.5%. As expected, opportunistic pathogens showed together the lowest relative IF contribution in all the subjects under study (from 5 to 10%).

Discussion
In these last years, 16S rRNA microarrays emerged as a sensitive and efficient way to screen complex bacterial communities. Here we describe and validate the HTF-Microbi.Array, a new phylogenetic DNA microarray designed for the high taxonomic level fingerprint of the human intestinal microbial community. The HTF-Microbi.Array is based on the LDR-UA approach, which is a fast and sensitive tool for the characterization of complex microbial communities with high sensitivity and specificity [25,26]. The use of this molecular technique allows overcoming the major limitations of DNA microarrays whose discriminative power is based on hybridization. In fact, a) optimization of the hybridiza-
tion conditions for each probe set is not required; b) problems due to the secondary structures of the target DNA are minimized, c) steric hindrances of differentially sized nucleic acid hybrids formed on the array after the hybridization are decreased [29]. The final probe set of the HTF-Microbi.Array allows a high taxonomic level fingerprint of the human intestinal microbiota, with a good coverage of the major and minor components, as well as some of the most important pathogens and opportunistic bacteria [30]. The LDR probes were designed by choosing DS oligonucleotides whose 3’end allowed the perfect discrimination of the target species from the non-target ones on the basis of our 16S rRNA sequence database. Definition of accurate and specific negative sets of gut microbiota sequences by ORMA tool [31] allowed the selection of maximally discriminative probe pairs. Probe specificity was confirmed on the entire known 16S rRNA gene sequences environment by the RDP Probe Match tool. This requirement is fundamental, since the primer set used for the PCR amplification was the “universal” 16S rRNA primer set designed by Edwards and co-workers [32].

The HTF-Microbi.Array recognized without ambiguity the 16S rRNA amplicons obtained from 28 members of the intestinal microbiota belonging to Bacteroides/Prevotella (B/P) blue, Clostridium cluster IV (Cl.IV) green, Clostridium cluster IX (Cl.IX) brown, Clostridium cluster XIVa (Cl.XIVa) dark brown. Lactobacillus, B. clausii, B. subtilis, Fusobacterium and Cyanobacteria are grouped as minor mutualistic symbionts (minor) indicated in yellow. Proteus, Yersinia and E. faecalis are grouped as opportunistic pathogens (opp) in red.

The HTF-Microbi.Array recognized without ambiguity the 16S rRNA amplicons obtained from 28 members of the intestinal microbiota belonging to Bacteroides/Prevotella (B/P) blue, Clostridium cluster IV (Cl.IV) green, Clostridium cluster IX (Cl.IX) brown, Clostridium cluster XIVa (Cl.XIVa) dark brown. Lactobacillus, B. clausii, B. subtilis, Fusobacterium and Cyanobacteria are grouped as minor mutualistic symbionts (minor) indicated in yellow. Proteus, Yersinia and E. faecalis are grouped as opportunistic pathogens (opp) in red.
is comparable to the values obtained by Rajilic-Stojanovic et al. [23] and Palmer et al. [21]. Thus the HTF-Microbi.Array shows the potentiality to sense low abundant species of the gastrointestinal microbiota, enabling the detection of the 16S rRNA of a peculiar target group present at a fractional abundance <0.1% in an artificial mixture.

The HTF-Microbi.Array was used in a pilot study to characterize the faecal microbiota of eight young adults. Faecal microbiota was chosen as DNA source since sample collection is not invasive, samples contain large amount of microbes, and, most important, it is representative of interpersonal differences in distal gut microbial ecology [33]. In order to have a good representation of the less abundant species of the intestinal microbial community, LDR reactions were performed starting from 50 fmol of PCR product. Cluster analysis of the presence-absence probes profiles enabled the identification of a reproducible high taxonomic level microbiota fingerprint for each subject. As expected, the intestinal microbial community of the voluntaries in the study resembled the typical fingerprint of healthy adults [28]. According to our data, the faecal microbiota of the enrolled subjects was dominated by major mutualistic symbionts. In fact, members of Bacteroidetes, Clostridium clusters IV, IX and XIVa were all represented in 100% of the subjects. On the other hand, minor mutualistic symbionts, such as Lactobacillaceae, B. subtilis et rel., Fusobacterium and Cyanobacteria, were detected in 55, 37, 50, and 63% of the subjects, respectively. Opportunistic pathogens, such as E. faecalis et rel., members of the Clostridium cluster I and II and Enterobacteriaceae, were represented only in 43, 25 and 12% of the subjects, respectively. Most importantly, enteropathogens such as, C. difficile, C. perfringens, E. faecium et rel., B. cereus et rel., and Campylobacter were never detected. A discrepancy between our data and the literature is the relatively low prevalence of the health promoting Bifidobacteriaceae in our samples (only 13% of samples). However, the low prevalence of bifidobacteria is a typical bias for several phylogenetic DNA microarrays [22,23]. Probably this is due to the intrinsic low efficiency of amplification of the bifidobacterial genome with universal primer sets for the 16S rRNA gene [8]. Surprisingly, a high prevalence was obtained for the minor mutualistic symbiont B. clausii et rel., 100% of samples, and the opportunistic pathogen Proteus, 50% of samples. For each subject the relative IF contributions of the probes were calculated, obtaining an approximate evaluation of the relative abundance of the principal microbial groups of the faecal microbiota. In general agreement with previous metagenomic studies [7-11] and SSU rRNA phylogenetic microarray investigations [22,23], mutualistic symbionts such as Bacteroidetes, Clostridium clusters IV, IX and XIVa largely dominated the faecal microbiota, contributing for the 65 to 80% of total microbiota, depending on the subject. Differently, with an overall contribution ranging from 10 to 30%, minor mutualistic symbionts such as B. clausii et rel., Bifidobacteriaceae, Lactobacillaceae, B. subtilis et rel., Fusobacterium, and Cyanobacteria were largely sub-dominant. Opportunistic pathogens represented only a small fraction of the intestinal microbiota. Even if subjects under study show a common trend when the ratio between the relative IF of major, minor and opportunistic components were considered, differences in the relative IF contribution of single probes were detectable and subject specific profiles were identified. For instance, subject n. 1 showed a higher relative fluorescence for probes targeting major mutualistic symbionts and a lower relative fluorescence for minor mutualistic symbionts and opportunistic pathogens than subjects n. 4 and 15. On the other hand subjects n. 15 and 17 were characterized by a lower ratio Bacteroidetes/Firmicutes with respect to all the other subjects. It is tempting to hypothesize that differences in relative IF contribution within samples could represent an approximation of differences in relative abundances of the targeted groups in the faecal microbiota. However, caution must be taken when microarray based methods for the relative quantification of bacterial groups in complex microbial communities are used. In fact, biases are introduced at several levels of the experimental procedure: DNA extraction and purification, PCR amplification of the 16S rRNA gene, and interspecies variation of the rRNA gene copy number [21].

Conclusion

The HTF-Microbi.Array has been revealed a fast and sensitive tool for the high taxonomic level fingerprint of the human intestinal microbiota in terms of presence/absence of the principal groups. Since the flexibility of the universal array platform allow the addition of new probe pairs without a further optimization of the hybridization conditions [25,26], the HTF-Microbi.Array can be easily implemented with the addition of new probe pairs targeting emerging microbial groups of the human intestinal microbiota, such as, for instance, the mucin degrading bacterium Akkermansia muciniphila [34]. The evaluation of the relative abundance of the target groups on the bases of the relative IF probes response still has some hindrances. However, considered all the possible biases (i.e. DNA extraction/purification, PCR, copy number variations, etc.) typical of the microarray technology, analysis of IFs from our LDR-UA platform can be useful in the estimation of the relative abundance of the targets groups within each sample. Focusing the phylogenetic resolution at division, order and cluster levels, the HTF-Microbi.Array results blind with respect to the inter-individual variability at the species level. Its potential to char-
acterize the high order taxonomic unbalances of the human intestinal microbiota associated with specific diseases will be assessed in further studies.

Methods

Recruitment

Eight healthy Italian individuals of 30 years old were enrolled for the study. None of the subjects had dietary restrictions except for antibiotics, probiotics and functional foods for at least 4 weeks prior to sampling. None of the selected subjects had a history of gastrointestinal disorders at the time of sampling. The study protocol was approved by the Ethical committee of Sant’Orsola-Malpighi Hospital (Bologna, Italy) and an informed consent was obtained from each enrolled subject. Faeces were collected for each subject and stored at -20°C.

Bacterial strains and culture conditions

The bifidobacterial strains used in this study were Bifidobacterium adolescentis ATCC15703, B. bifidum DSM20456, B. breve DSM20091, B. longum ATCC15707. The Lactobacillus strains were Lactobacillus plantarum DSM21074, L. casei DSM20011, L. rhamnusus DSM20021, L. salivarius SV2 (strain from our collection), L. delbrueckii DSM 20314, L. gasseri DSM20243, L. reuteri DSM20016, L. pentosus DSM20134, L. acidophilus DSM20079. All bifidobacteria and Lactobacillus strains were grown on De Man-Rogosa-Sharpe (MRS) broth with cysteine (0.5 g/l) at 37°C under an anaerobic atmosphere (AnaeroCult, Merck, Darmstadt, Germany). Escherichia coli ATCC111015 was cultivated at 37°C aerobically on TY-broth. Salmonella cholerasuis typhimurium and Yersinia enterocolitica-type, kindly provided by A. Essig, Dept. of Medical Microbiology, University of Ulm, Germany, were cultivated aerobically at 30°C on BHI-broth.

Target selection and consensus extraction

A database of 16S rRNA sequences was created by integration of the 16S rRNA database of the ARB Project (release February, 2005) [http://www.arb-home.de; [35]] with the database of the Ribosomal Database Project (RDP; release September, 2007) [http://rdp.cme.msu.edu/; [36,37]]. A phylogenetic tree was obtained in the ARB software, by using the neighbour-joining algorithm for the sequence alignment. The tree was used for the rational selection of phylogenetically related groups of bacteria belonging to the human intestinal microbiota which correspond to nodes of the phylogenetic tree (Additional file 1). Group specific consensus sequences were extracted, with a cut-off of 75% for base calling. Nucleotides which occurred at lower frequencies were replaced by the appropriate IUPAC ambiguity code.

Probe design

Multiple alignment step of the selected sequences was performed in ClustalW [38]. Since the taxonomic classification of the 30 groups selected for the probe design varied from species to phylum level, careful grouping of the sequences was performed for the multiple alignment step: (a) for higher level probes, only family/phylum consensus sequences were used as a negative set for probe design; (b) for genus/species level probes, only sequences belonging to other families/phyla were selected. All the LDR probe pairs were designed using ORMA [31]. Both DS and CP were required to be between 25 and 60 bases pair, with a Tm of 68 ± 1°C, and with maximum 4 degenerated bases. In-silico check versus a publicly available database (i.e.: RDP) was then performed for assessing probe pair specificity.

DNA extraction

Total DNA was extracted from 10^9 bacterial cells by using the DNeasy Tissue Kit 50 (Qiagen, Düsseldorf, Germany) following the manufacturer instructions. Bacterial DNA was also extracted from lyophilized bacterial cells of the following DSMZ (Braunschweig, Germany) collection strains: Clostridium leptum DSM73, Ruminococcus albus DSM20455, Eubacterium siraeum DSM15700, C. viride DSM6836, Megasphaera micruniciformis DSM17226, Bacillus clausii DSM2515, B. subtilis DSM704, B. cereus DSM21, and Proteus mirabilis DSM4479. Lyophilized bacterial cells were suspended in 1 ml of lysis buffer (500 mM NaCl, 50 mM Tris-HCl pH 8, 50 mM EDTA, 4% SDS) and DNA extraction was carried out by employing the same procedure used for the extraction of genomic DNA from faecal samples, according to the following procedure. Total DNA from faecal material was extracted using QIAamp DNA Stool Min Kit (Qiagen) with a modified protocol. 250 mg of faeces were suspended in 1 ml of lysis buffer. Four 3 mm glass beads and 0.5 g of 0.1 mm zirconia beads were added, and the samples were treated in FastPrep (MP Biomedical, Irvine, CA, USA) at 5.5 ms for 3 min. Samples were heated at 95°C for 15 minutes, then centrifuged for 5 min at full speed to pellet stool particles. Supernatants were collected and 260 μl of 10 M ammonium acetate were added, followed by incubation in ice for 5 min and centrifugation at full speed for 10 min. One volume of isopropanol was added to each supernatant and incubated in ice for 30 min. The precipitated nucleic acids were collected by centrifugation for 15 min at full speed and washed with 70% ethanol. Pellets were resuspended in 100 μl of TE buffer and treated with 2 μl of DNase-free RNase (10 mg/ml) at 37°C for 15 min. Protein removal by Proteinase K treatment and DNA purification with QIAamp Mini Spin columns were performed following the kit protocol. 200 μl of TE buffer were used for DNA elution. Final DNA
concentration was determined by using NanoDrop ND-1000 (NanoDrop Technologies, Wilmington, DE). The bacterial DNA from the following 11 ATCC strains was directly obtained from the ATCC: *Bacteroides fragilis* ATCC25285, *B. thetaiotaomicron* ATCC29148, *Prevotella melaninogenica* ATCC25845, *Veillonella parvula* ATCC10790, *C. difficile* ATCCBAA1382, *C. acetobutylicum* ATCC824, *C. perfringens* ATCC13124, *Enterococcus faecalis* ATCC700802, *E. faecium* ATCC51559, *Campylobacter jejuni* ATCC33292, *R. productus* 23340.

**Polymerase Chain Reaction (PCR)**

All the oligonucleotide primers and probe pairs were synthesized by Thermo Electron (Ulm, Germany). PCR amplifications were performed with Biometra Thermal Cycler II and Biometra Thermal Cycler T Gradient (Biometra, Germany). PCR products were purified by using a Wizard SV gel and PCR clean-up System purification kit (Promega Italia, Milan, Italy), according to the manufacturer’s instructions, eluted in 20 μl of sterile water, and quantified with the DNA 7500 LabChip Assay kit and BioAnalyzer 2100 (Agilent Technologies, Palo Alto, CA, USA). 16S rRNA was amplified using universal forward primer 16S27F (5'-AGAGTTTGTCTGCTGGGCTCAG-3') and reverse primer r1492 (5'-TACGGYTACCTTGTTACGACTT-3'), following the protocol described in Castiglioni et al. [25] except for using 50 ng of starting DNA and 0.5 U of DNAzyme DNA polymerase II (Finnzymes, Espoo, Finland).

**LDR/Universal Array approach**

Phenylen-diisothiocyanate (PDITC) activated chitosan glass slides were used as surfaces for the preparation of universal arrays [39], comprising a total of 49 Zip-codes. Hybridization controls (cZip 66 oligonucleotide, complementary to zip 66, 5'-Cy3-GTTACCGCTGGTGCTGCGCCGCT-3') were used to locate the submatrices during the scanning. The entire experimental procedure for both the chemical treatment and the spotting is described in detail in Consolandi et al. [40]. An overview of the Universal Array layout and ZipCodes is provided as Additional file 6. Ligase Detection Reaction and hybridization of the products on the universal arrays were performed according to the protocol described in Castiglioni et al. [25] except for the probe annealing temperature, set at 60°C.

The LDRs were carried out in a final volume of 20 μl with different quantities of purified PCR products: a) all LDRs for specificity tests were performed on 50 fmol of initial PCR product, for having no issues related to target; b) sensitivity tests were performed with decreasing PCR product concentration from 75 to 0.7 fmol; c) relative abundance tests were performed on 1 fmol *E. coli* PCR amplicon, mixed with human genomic DNA extracted from whole blood, at decreasing concentrations, from 4%, down to 0.02%; d) LDR experiments on the eight faecal samples were performed on 50 fmol PCR product.

**Data analysis**

All arrays were scanned with ScanArray 5000 scanner (Perkin Elmer Life Sciences, Boston, MA, USA), at 10 μm resolution. In the experiments, the fluorescent images were obtained with different acquisition parameters on both laser power and photo-multiplier gain, in order to avoid saturation. IF were quantitated by ScanArray Express 3.0 software, using the "Adaptive circle" option, letting diameters vary from 60 to 300 μm. No normalization procedures on the IFs have been performed. To assess whether a probe pair was significantly above the background (i.e. was 'present' or not), we performed a one-sided t-test (α = 0.01). The criteria was relaxed to α = 0.05 for sensitivity tests. The null distribution was set as the population of "Blank" spots (e.g. with no oligonucleotide spotted, n = 6). Two times the standard deviation of pixel intensities of the same spots was added to obtain a conservative estimate. For each zip-code, we considered the population of the IFs of all the replicates (n = 4) and tested it for being significantly above the null-distribution ($H_0: \mu_{test} = \mu_{null}; H_1: \mu_{test} > \mu_{null}$). In case one replicate in the test population was below 2.5 times the distribution mean, this was considered an outlier and was discarded from the analyses. We calculated the ratio between the signal intensities of the specific probes on the blank intensity (SNR), and the ratio between all the other probes and the blank intensity (SNR$^{nr}$).

**Clustering**

Hierarchical clustering of HTF-Microbi.Array profiles was carried out using the statistical software R [http://www.r-project.org]. The Euclidean distance among sample profiles was calculated and Ward's method was used for agglomeration.

**Additional material**

- **Additional file 1** HTF-Microbi.Array target groups. Phylogenetically related groups target of the HTF-Microbi.Array.
- **Additional file 2** HTF-Microbi.Array probe list. Table of the 30 designed probe pairs. Sequences (5’ -> 3’) for both DS and CP are reported, as well as major thermodynamic parameters (melting temperature, length, number of degenerated bases).
- **Additional file 3** Specificity tests of the HTF-Microbi.Array. Raw data of the specificity tests of the HTF-Microbi.Array. Each column represents a different sample, whose identification is reported as its label. On the left, the ZipCode, the probe name and ID are reported. "Type" is a numeric flag used for the classification of the probes: 1 is the hybridization control, 2 is the ligation control, 3 indicates the HTF-Microbi.Array probes, 4 are the unused ZipCodes and 5 is the Blank. "Numeric ID" is given to the probes according to their "type" and "Oligo ID" values.
Additional file 4 Sensitivity tests of the HTF-Microbi.Array
Raw data of the sensitivity tests on the HTF-Microbi.Array. The workbook has two spreadsheets: "Artificial mix data", reporting the results of the serial dilutions of the 6 bacterial DNA mix (B. cereus, E. coli, S. aureus, Pre-votella, Y. enterocolitica), with concentrations ranging from 50 to 0.7 fmol. "Absolute sensitivity E. coli" spreadsheet reports the results of the tests on low quantities of E. coli 16S amplicon in increasing amounts of human genomic DNA. The file is structured as described above for Additional file 3.

Additional file 5 Tests of the HTF-Microbi.Array on faecal samples
Raw data for the experimental characterization of the faecal microbiota of eight healthy young adults. Patient ID and replicate number are reported as the column headers. The file is structured as described above for Additional file 3.

Additional file 6 Universal array scheme
Graphical representation of the Universal Array platform. Each array has 8 identical subarrays (A), which can be addressed independently. Each subarray is made by 208 spots, with quadruplicates of each ZipCode (B); hybridization and ligation controls and Blanks are repeated 8, 6 and 6 times, respectively; the figure highlights in gray the ZipCodes actually associated to probe pairs used in the HTF-Microbi.Array. Sequences (5' - 3' oriented) and numbers of the ZipCodes are reported in (C).

Authors’ contributions
MC, CC, MS, and EB performed the study design, analysis and interpretation of the data and the writing of the paper. BC and BV participated in the design of the study. GDB and PB coordinated the study. All authors read and approved the manuscript.

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Author Details
1Department of Pharmaceutical Science, University of Bologna, Italy, 2Institute of Agricultural Biology and Biotechnology - Italian National Research Council, Milan, Italy and 3Institute of Biomedical Technologies - Italian National Research Council, Milan, Italy

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