Microbial Community Structure of Three Traditional Zambian Fermented Products: Mabisi, Chibwantu and Munkoyo

Sijmen E. Schoustra1,2,3, Chitundu Kasase3, Cristian Toarta2, Rees Kassen2, Alexandre J. Poulain2*

1 Laboratory of Genetics, Wageningen University, Wageningen, Netherlands, 2 Department of Biology, University of Ottawa, Ottawa, Canada, 3 Departments of Biological Sciences and Food Science and Technology, University of Zambia, Lusaka, Zambia

Abstract

Around the world, raw materials are converted into fermented food products through microbial and enzymatic activity. Products are typically produced using a process known as batch culture, where small volumes of an old culture are used to initiate a fresh culture. Repeated over many years, and provided samples are not shared among producers, batch culture techniques allow for the natural evolution of independent microbial ecosystems. While these products form an important part of the diets of many people because of their nutritional, organoleptic and food safety properties, for many traditional African fermented products the microbial communities responsible for fermentation are largely unknown. Here we describe the microbial composition of three traditional fermented non-alcoholic beverages that are widely consumed across Zambia: the milk based product Mabisi and the cereal based products Munkoyo and Chibwantu. Using culture and non-culture based techniques, we found that six to eight lactic acid bacteria predominate in all products. We then used this data to investigate in more detail the factors affecting community structure. We found that products made from similar raw materials do not harbor microbial communities that are more similar to each other than those made from different raw materials. We also found that samples from the same product taken at the same location were as different from each other in terms of microbial community structure and composition, as those from geographically very distant locations. These results suggest that microbial community structure in these products is neither a simple consequence of the raw materials used, nor the particular suite of microbes available in the environment but that anthropogenic variables (e.g., competition among sellers or organoleptic preferences by different tribes) are important in shaping the microbial community structures.

Introduction

Fermented foods are a significant part of the daily diet of many people in Africa and around the world [1]. During production of a fermented product, microorganisms transform raw material into a product with increased value, generally by extending the shelf life of the raw materials, by increasing the nutritional value of the product and/or by improving the product’s organoleptic attributes [2,3]. In most cases, starter culture is added by transferring some of an old batch to fresh raw materials, a process referred to as back-slopping by producers. Back-slopping resembles very closely the back-slopping by producers. Back-slopping resembles very closely the batch culture transfer procedure used by microbiologists to study the evolution of microbial populations and communities in the laboratory. When repeated over numerous cycles [1] this process can result in a co-adapted, evolutionarily stable microbial community [4].

The diversity of fermented products found in most African countries is influenced by the available local raw materials and ingredients as well as, to a large extent, by indigenous knowledge and practices. For several African fermented products, the microbial flora has been characterized, critical steps in the process have been identified, and production industrialized. In many cases, the microbes responsible for fermentation are lactic acid bacteria and yeasts that are added to raw materials as a starter culture. These microbes drive the fermentation and hence determine the type of final product and its properties. However, for most African products details on the fermentation process are largely unknown [3,5,6,7,8].

In Zambia numerous endogenous fermented products are produced, of which Munkoyo, Chibwantu and Mabisi are three important examples of non-alcoholic beverages [5,6]. These products are consumed on a daily basis by many people and have a great socio-economic and cultural importance in Zambia and Southern Africa. The main motivation for local people to produce these products is their appealing taste, improved nutritional value and prolonged shelf life. In addition, local people claim that the products offer health benefits, such as the prevention and cure of diarrhea. The products are produced at home and almost exclusively by women who sell their product at local markets, providing them with an additional source of income.

The microbial ecology of these three traditional fermented beverages is unknown and forms the focus of this study. We
analysed the community structure for products of the same denomination and the same starting raw material (milk for Mabisi, cereal for Chibwantu and Munkoyo) found at different geographical locations and prepared by different tribes. To analyse communities, we used both culture based and non-culture based techniques and this study allows a comparison of the efficiency of these two complementary approaches for the microbial analysis of these products. Our study had two main objectives: (1) to characterise the microbial community structures of the products in an effort to support and develop traditional knowledge, with an eye to eventually facilitating potential process optimisation and economic development, and (2) to obtain some basic understanding of two possible factors governing microbial community structure in these fermented products. More specifically, we asked if starting materials or sampling location is important in structuring the microbial communities within these products.

Methods

Preparation and Description of the Products

Munkoyo and Chibwantu are non-alcoholic beverages based on cereal raw materials and are produced in a similar way, although the end products have slightly different tastes and organoleptic properties (Figure 1A). Munkoyo is made in the Copperbelt province and northern parts of Zambia; Chibwantu is produced in the Central and Southern areas including those around the capital Lusaka. Both products are made from maize (with sometimes the addition of millet and/or sorghum). Roots of certain plants (Eminia, Rynchosia and Vigna species; [6]) are added after soaking of the powdered grains and are thought to provide amylolytic enzymes, which make available starch in the cereals by degrading it into maltose (80%), maltotriose (17%) and glucose (3%). The roots are from shrubs that grow as under storey plants in the Miombo Woodland. The roots are tuber-like and fibrous (3%). The roots are from shrubs that grow as under storey plants in the Miombo Woodland. The roots are tuber-like and fibrous (3%). The roots are from shrubs that grow as under storey plants in the Miombo Woodland. The roots are tuber-like and fibrous (3%). The roots are from shrubs that grow as under storey plants in the Miombo Woodland. The roots are tuber-like and fibrous (3%). The roots are from shrubs that grow as under storey plants in the Miombo Woodland. The roots are tuber-like and fibrous (3%). The roots are from shrubs that grow as under storey plants in the Miombo Woodland. The roots are tuber-like and fibrous (3%). The roots are from shrubs that grow as under storey plants in the Miombo Woodland. The roots are tuber-like and fibrous (3%).

The products are also major beverages consumed during special ceremonies. Over the last few years, variants of Munkoyo and Chibwantu (Maheu) have become commercially available via an alternative production process that involves only the addition of enzymes and artificial flavourings. However, no fermentation by live microbial cultures is involved and many people still prefer the traditional versions (Munkoyo and Chibwantu). The end products have slightly different tastes and organoleptic properties.

Chibwantu is produced in the Copperbelt province and northern parts of Zambia; Chibwantu is produced in the Central and Southern areas including those around the capital Lusaka. Both products are made from maize (with sometimes the addition of millet and/or sorghum). Roots of certain plants (Eminia, Rynchosia and Vigna species; [6]) are added after soaking of the powdered grains and are thought to provide amylolytic enzymes, which make available starch in the cereals by degrading it into maltose (80%), maltotriose (17%) and glucose (3%). The roots are from shrubs that grow as under storey plants in the Miombo Woodland. The roots are tuber-like and fibrous (3%). The roots are from shrubs that grow as under storey plants in the Miombo Woodland. The roots are tuber-like and fibrous (3%). The roots are from shrubs that grow as under storey plants in the Miombo Woodland. The roots are tuber-like and fibrous (3%). The roots are from shrubs that grow as under storey plants in the Miombo Woodland. The roots are tuber-like and fibrous (3%). The roots are from shrubs that grow as under storey plants in the Miombo Woodland. The roots are tuber-like and fibrous (3%). The roots are from shrubs that grow as under storey plants in the Miombo Woodland. The roots are tuber-like and fibrous (3%). The roots are from shrubs that grow as under storey plants in the Miombo Woodland. The roots are tuber-like and fibrous (3%).

Sampling

Samples of approximately 250 mL were purchased from producers at local markets (Table 1; Figure 1C). pH of each sample was measured at the sampling site. Samples (2×50 mL and 2×15 mL of the original 250 mL) were then transferred to the laboratory and stored at −20°C and −80°C. Prior to analysis, samples were well mixed by vortexing for at least 45 seconds. Thirty-seven samples were collected in total.

Culture Based Techniques

Samples were serially diluted in Minimal Salts Medium (Na2HPO4 6.7 g, KH2PO4 3 g, NaCl 0.5 g, NH4Cl 1.0 g, 1000 mL dH2O). Dilutions were plated on the following solid growth media: Plate Count Agar (PCA; 0.3% peptone, 0.25% yeast extract, 0.1% glucose, 1.5% agar, pH 7) incubated at 28°C both aerobically and non-aerobically for total aerobic and facultative anaerobic counts; PCA plates were incubated aerobiically at 42°C for thermophiles; M17 plates (Oxoid) were incubated at 28°C aerobiically for lactic acid bacteria; MRS plates (Oxoid) were incubated at 28°C anaerobiically for facultative anaerobic lactic acid bacteria; YM plates were (0.3% yeast extract, 0.5% peptone, 1% agar, 0.5% malt extract, 1% glucose) at 28°C aerobiically for yeast and Violet Bile agar (VB; Oxoid) grown at 28°C for Enterococci. After 4 days of incubation, plates were examined and colonies were counted. From each medium, at least eight colonies were picked capturing all different colony morphologies (if any) and streaked on fresh solid medium for purification. As a result, we collected 30 to 42 bacterial isolates from each sample. These isolates were identified by sequencing part of the V1 to V4 region of the 16S ribosomal DNA gene (see below under PCR amplification).

DNA Extraction

We designed and optimised DNA extraction procedures for each product. For the cereal based products (Chibwantu and Munkoyo), the protocol for DNA extraction was adapted from Abriouel et al. 2006 and Ampe et al. 1999 [9,10]. First, we eliminated large particles from 1 gram of product. Samples were spun down at high speed, supernatant was removed and we kept the pellet. We added 500 µl TESL (25 mM Tris, 10 mM EDTA, 20% sucrose, 20 mg/ml lysozyme) and 10 µl metanolysis solution (in water at 1 U/µl), followed by incubation at 37°C for 60 minutes with slight shaking. We added 500 µl GES reagent (5 M guanidinium thiocyanate, 100 mM EDTA, 0.5% sarcosyl), cooled on ice for 5 minutes and added 250 µl of cold ammonium acetate solution (7.5 M), followed by gentle mixing. The mixture was held on ice for 10 minutes, spun down and the supernatant removed. We purified the samples by mixing with chloroform-2-pentanol mix (chloroform and 2- pentanol 24:1) by adding 1:1 to the supernatant; we spun down and kept the supernatant. We performed a phenol-chloroform purification by adding equal volume of phenol (i.e., tris-saturated phenol-chloroform-isooamy-lethanol at a ratio of 24:25:1), vortexed for a few seconds, spun for 2 minutes at 12000 rpm 4°C and transferred the supernatant to a fresh tube. We then added equal volume of chloroform to the supernatant, vortexed for a few seconds, spun 2 minutes at 12000 rpm and 4°C and transferred the supernatant to a fresh tube. We then added 2.5 volumes 100% ethanol, vortexed and precipitated DNA at −20°C overnight. Subsequently, samples were spun for 20 minutes at 12000 rpm and 4°C; the supernatant was removed by aspiration. We washed the DNA by adding 1 mL cold 70% ethanol, spun for 10 minutes at 12000 rpm and 4°C; the supernatant was removed by aspiration and the DNA pellet was air-dried for 10 minutes at room temperature. Finally, the DNA was dissolved in 10 mM Tris treated with RNase (10 mM Tris, bring to pH 8.0 with HCl, 1 mM EDTA, RNase 20 µl/ml).
Figure 1. Preparation of the fermented products and sampling locations. A). Process flow for the preparation of Monkoyo. Chibwantu is produced in the same way. If the product is allowed to be fermented for an additional 5 days an alcoholic drink develops that can be kept for prolonged periods of time. B). Process flow of Mabisi. C). Map of Zambia indicating sampling locations (see Table 1). D). Picture of the fermentation vessel (Calabash). doi:10.1371/journal.pone.0063948.g001
We added 400 μg/ml protenase E and incubated for 60 minutes at 37°C. The protocol was performed as follows (adapted from Ercolini et al.): we transferred the supernatant containing the DNA to a fresh 1.5 ml microcentrifuge tube and centrifuged the sample at 13,000 × g for 10 minutes. We performed this PCR step either for selected purified single isolates, or on the DNA extracted from an entire sample to generate clone libraries. GoTaq and Go buffer supplied by Promega were used for the reaction following the manufacturer’s instructions. For each 25 μl reaction, 2 μl of a 25 mM MgCl₂ solution was added to the reaction buffer. PCR conditions were as follows: denaturation at 94°C for 2 minutes; then 33 cycles of denaturing at 94°C for 30 sec, annealing at 55°C for 30 sec, elongation at 72°C for 90 sec; followed by a final elongation step at 72°C for 10 minutes.

### Table 1. A total of 37 samples were collected of three product-types; Chibwantu, Mabisi and Munkoyo in various towns across Zambia (Figure 1C).

| SampleID | Product type | pH  | Town      |
|----------|--------------|-----|-----------|
| Chibw-1  | Chibwantu    | 4.2 | Mumbwa    |
| Chibw-1B | Chibwantu    | 4.2 | Mumbwa    |
| Chibw-2  | Chibwantu    | 4.2 | Mumbwa    |
| Chibw-3  | Chibwantu    | 4.5 | Mumbwa    |
| Chibw-4  | Chibwantu    | 4.3 | Mumbwa    |
| Chibw-5  | Chibwantu    | 4.4 | Mumbwa    |
| Chibw-6  | Chibwantu    | 4.0 | Monze     |
| Chibw-7  | Chibwantu    | 3.8 | Choma     |
| Chibw-8  | Chibwantu    | 3.8 | Choma     |
| Chibw-9  | Chibwantu    | 4.0 | Lusaka    |
| Mabisi-1 | Mabisi       | 4.5 | Mumbwa    |
| Mabisi-2 | Mabisi       | 4.4 | Mumbwa    |
| Mabisi-3 | Mabisi       | 4.5 | Mumbwa    |
| Mabisi-1B| Mabisi       | 4.4 | Mumbwa    |
| Mabisi-2B| Mabisi       | 4.5 | Mumbwa    |
| Mabisi-3B| Mabisi       | 4.3 | Mumbwa    |
| Mabisi-5 | Mabisi       | 4.2 | Monze     |
| Mabisi-6 | Mabisi       | 4.0 | Zimba     |
| Mabisi-6B| Mabisi       | 4.0 | Zimba     |
| Mabisi-7 | Mabisi       | 4.0 | Choma     |
| Mabisi-8 | Mabisi       | 4.0 | Choma     |
| Mabisi-9 | Mabisi       | 4.0 | Choma     |
| Mabisi-10| Mabisi       | 4.0 | Choma     |
| Munkoyo-1| Munkoyo      | 3.5 | Mufulira  |
| Munkoyo-2| Munkoyo      | 3.5 | Mufulira  |
| Munkoyo-4| Munkoyo      | 3.5 | Mufulira  |
| Munkoyo-5| Munkoyo      | 3.5 | Mufulira  |
| Munkoyo-6| Munkoyo      | 3.5 | Mufulira  |
| Munkoyo-7| Munkoyo      | 3.5 | Mufulira  |
| Munkoyo-8| Munkoyo      | 3.5 | Mufulira  |
| Munkoyo-9| Munkoyo      | 3.6 | Choma     |
| Unza-1   | Munkoyo      | 4.0 | Lusaka    |
| Unza-2   | Munkoyo      | 3.8 | Lusaka    |
| Unza-3   | Munkoyo      | 3.8 | Lusaka    |
| Unza-4   | Munkoyo      | 3.8 | Lusaka    |
| Unza-5   | Munkoyo      | 4.0 | Lusaka    |

Immediately after sampling, the pH of the sample was measured.

For the milk-based product (Mabisi) the DNA extraction protocol was performed as follows (adapted from Ercole et al. 2001, [11]): we started with 1 ml of product in a 1.5 ml microcentrifuge tube and centrifuged the sample at 13,000 × g for 2 minutes to pellet the cells and removed the supernatant. We resuspended the cells in a solution containing 64 μl of 1 M ammonium acetate (5 M) and cooled on ice for 15 minutes before being spun down at 13,000 × g for 10 minutes. We transferred the supernatant containing the DNA to a fresh 1.5 ml microcentrifuge tube and performed a phenol-chloroform DNA purification as described above.

### PCR Amplification

We used primers 27F (forward primer 5’AGAGTTTGATCMTGGCTCAG’3) and 907R (reverse primer 5’CCGCTAACATTTGAGCTACGG’3) to amplify an 880 bp fragment of the V1 to V4 hypervariable regions on the 16S rDNA gene. PCR products using a combination of EcoRI and AluI for the Chibwantu and Munkoyo libraries and EcoRI and HindIII for the Mabisi libraries. In a total volume of 25 μl, 1 μl of PCR product was digested overnight at room temperature with at least 500 units of each enzyme using appropriate reaction buffer as per the manufacturer’s instructions (New England BioLabs). Restriction patterns were separated on 1.75% agarose gels and stained with ethidium bromide. Each different restriction pattern was defined using MOTHUR [13].

### Cloning and Sequencing of 16S rRNA Gene PCR Products

PCR products were sequenced by gel electrophoresis (2% agarose). The 880 bp 16S rRNA gene amplicons were extracted and purified from the gel and cloned into vector pSC-A-amp/kan, using a Strataclone PCR cloning kit (Stratagene, Agilent Technologies) according to the manufacturer’s instructions. Eight libraries were created (three for Chibwantu, two for Munkoyo and three for Mabisi). Plasmid DNA was isolated from 96 clones from each library using a PureLink Promega Wizard Magnesium plasmid purification system and underwent ARDRA (see below). Selected clones containing the 880 bp insert were sequenced using the M13f primer (see below). All sequences have been submitted to GenBank, with accession numbers JQ807985 to JQ808112.

### ARDRA

For amplified ribosomal DNA restriction analysis (ARDRA), 48 randomly picked colonies per 16S rDNA gene library were each grown overnight at 37°C in LB containing ampicillin. Plasmid DNA extraction was performed using a Promega Wizard Magnesium plasmid purification system and the insert amplified using primers M13f and M13r. Double restriction digests were made of the PCR products using a combination of EcoRI and AluI for the Chibwantu and Munkoyo libraries and EcoRI and HindIII for the Mabisi libraries. In a total volume of 25 μl, 1 μl of PCR product was digested overnight at room temperature with at least 500 units of each enzyme using appropriate reaction buffer as per the manufacturer’s instructions (New England BioLabs). Restriction patterns were separated on 1.75% agarose gels and stained with ethidium bromide. Each different restriction pattern was defined as an operational taxonomic unit (OTU) (Figure S2). The distribution of OTUs in each library was determined. Two to three representatives of each OTU were sequenced using primers M13f and compared to those in the RDP database [12] or GenBank database using the BLAST algorithm. Diversity indices (Chao1 index, Shannon index, coverage, rarefaction) were computed using MOTHUR [13].

### Phylogenies and Sequences Analyses

Sequencing was performed by Genome Quebec at the sequencing facility of McGill University in Montreal (Canada). All 16S rDNA gene sequences generated were initially scanned for vectors using the PIPELINE function in the Ribosomal Database.
Microbial Characterization of Zambian Products

Project (RDP) [12] and manually inspected. Primers from 16S rRNA gene sequences were removed using Geneious software package primer annotation tool. Chimeric sequences were identified using Mallard, confirmed using Pintail and were discarded [14]. The 16S rRNA sequences were aligned with the latest 16S rRNA gene core alignment downloaded from the SILVA database using MOTHUR [13]. The model of evolution most appropriate to the dataset was determined using jModelTest v.0.1.1 [15]; phylogenies were constructed using FastTree [16] using the appropriate model of evolution and graphically represented using FigTree.

Microbial community structures for each product were compared based on phylogenetic information using the UniFrac metric [17]. For those samples that showed significant difference in the community composition, we performed Jackknife cluster analyses. Using UniFrac, we computed a (phylogenetic) similarity matrix based on clone library [17], using one clone library (we arbitrarily chose chibwantu1) as a scaling reference. We analysed how phylogenetic similarity correlates with geographical distance through linear regression (Figure S3).

Results

We collected a total of 37 samples of fermented products from Zambia (Table 1). We characterised the microbial community structures of 3 Zambian fermented products, Mabisi (milk based), Munkoyo and Chibwantu (cereal based) using culture and non-culture based techniques. The pH of the products ranged from 4.0 to 4.5 for Mabisi and Chibwantu and from 3.5 to 4.0 for Munkoyo.

Characterisation of Isolates and Microbial Communities

For the culture-based analysis we used 5 different growth media (see Methods) which revealed that total viable counts are on the order of $10^7$ cfu/g of product (Figure 2), with highest densities found in Mabisi and lowest in Munkoyo (ANOVA: $F_{2,35} = 6.35$, $p = 0.005$). As can be seen from the counts on M17 and MRS agar, facultative anaerobic lactic acid bacteria dominate the culturable components of the communities. This result is consistent with the low pH of the products (Table 1) and suggests that the fermentation vessel is oxygen limited (Figure 1D), providing an environment conducive to acid formation. We chose chibwantu1 as a scaling reference. We analysed how phylogenetic similarity correlates with geographical distance through linear regression (Figure S3).

We constructed clone libraries for 8 product samples (3 for Mabisi, 3 for Chibwantu and 2 for Munkoyo). Table 3 shows the number of distinct RFLP patterns and the abundance within each pattern. We selected up to 3 clones per RFLP type (11 to 15 clones per library) for sequencing. The identity of clones per observed RFLP pattern is in Table 3 and their phylogenetic relationship is presented in Figure 3 together with several reference strains from the NCBI database. Members of the Lactobacillus and Weisella genera dominated Chibwantu products; Mabisi products were dominated by members of the Lactococcus, Lactobacillus and Streptococcus genera and Munkoyo products were dominated by members of the Weisella and Lactobacillus genera. A few gram-negative genera were found, namely, Acinetobacter, Chryseobacterium, Actibacter and Gluconobacter (Table S1, Table 3 and Figure 3). In almost all cases, only one to four key players had abundances greater than 15%. Table S1 provides full details of all sequences we obtained.

We compared the community composition obtained through culture based and non-culture based approaches using the UniFrac significance metric for the 3 product samples (chibwantu8, mabisi10 and munkoyo5). Results showed that the microbial communities identified using culture-based or non culture-based approaches were not different from one another for Mabisi (UniFrac significance; $p = 0.13$); culture-based and non culture-based approaches yielded different community structures for Chibwantu and Munkoyo (UniFrac significance; $p<0.01$ in both cases).

Finally, our results revealed that we adequately sampled each of the libraries constructed for each of the products (Table 4). In all cases, using 48 clones was sufficient to capture the variation present. This is supported by Good’s coverage estimates varying from 93% to 98%, by the low values obtained by the Chao1 richness estimators and by the analyses of the rarefaction curves (Figure S1). Overall diversity, estimated by Shannon indices, was low ranging from 0.63 to 0.77 for Munkoyo to 0.39 to 2.20 for Chibwantu. Mabisi exhibited intermediate values ranging from 1.36 to 1.53 (Table 4).

Microbial Community Structure Comparisons

We used UniFrac tools to compare microbial community structures of the various fermented products investigated using phylogenetic information [16]. When compared using the UniFrac significance metric, community structures of Munkoyo, Mabisi and Chibwantu were significantly different from each other (Table 5A). The results of the cluster analysis (Figure 3 inset) revealed that in all cases, Munkoyo clustered away from Chibwantu and Mabisi, suggesting that the communities in Chibwantu and Mabisi are more similar to each other than the communities in Munkoyo. A Jackknife analysis provides further support for this result: Munkoyo clustered away from Chibwantu and Mabisi 100% of the time. Due to lack of statistical power, Mabisi and Chibwantu differences could only be retrieved 15% of the time with the Jackknife analysis.

We further analysed libraries of the same product type but from different markets or vendors and found that most microbial communities were significantly different from one another except for Mabisi (Table 5 B and C). It might be expected that, for a given product, microbial communities collected in close proximity would be more similar to each other than those from more distant locations. This was apparently not the case, at least for Chibwantu and Mabisi. Libraries of samples taken at geographically proximal locations for these products did not group together. A cluster analysis on all product samples, pooled based on their sampling location, also found that samples collected in proximity to each
other did not group together. For example, the distance between the towns of Choma and Monze is about 50 km and that between Choma and Mumbwa is 350 km (Figure 1D) but the microbial community of samples collected in Choma and Monze are apparently the most different in our analysis (Figure 4; Table 5D).

Furthermore there is no significant negative relationship between the geographical distance at which the samples were collected and the similarity of the microbial community composition, as would be expected if microbial community similarity declines with distance. Using one library (Chibwantu1) as a reference and

**Table 2.** List of strains isolated from product samples using culture based techniques.

| Product | Isolate | BLAST identity                  | Abundance (per g of product) |
|---------|---------|---------------------------------|------------------------------|
| Chibwantu | Chibwantu03 | *Bacillus safensis*                  | 10⁵                           |
|         | Chibwantu07 | *Gluconobacter oxydans*            | 10⁵                           |
|         | Chibwantu08 | *Leuconostoc garlicum*             | 10⁵                           |
|         | Chibwantu10 | *Bacillus subtilis*                | 10⁵                           |
|         | Chibwantu11 | *Paenibacillus motobuensis*        | 10⁴                           |
| Mabisi  | Mabisi28 | *Lactobacillus brevis*             | 10⁷                           |
|         | Mabisi19 | *Lactobacillus plantarum*          | 10⁷                           |
|         | Mabisi09 | *Leuconostoc garlicum*             | 10⁵                           |
|         | Mabisi11 | *Enterococcus durans*              | 10⁵                           |
|         | Mabisi15 | *Lysinibacillus sphaericus*        | 10⁴                           |
| Munkoyo | Munkoyo09 | *Lactobacillus fermentum*          | 10⁵                           |
|         | Munkoyo13 | *Lactobacillus plantarum*          | 10⁵                           |
|         | Munkoyo15 | *Bacillus lecheniformis*           | 10⁴                           |
|         | Munkoyo11 | *Lactobacillus brevis*             | 10⁵                           |

Based on the dilution at which the strain was isolated, the abundance was estimated. See the Table S1 for full details on the sequences that we obtained to identify each strain.

doi:10.1371/journal.pone.0063948.t002
Figure 3. Phylogenetic tree. Maximum likelihood tree of bacterial 16S rRNA gene sequences. Triangle heights are proportional to the number of sequences. Numbers adjacent to each node represent aLRT statistics (SH-like supports). Scale bar shows branch lengths (expected number of substitutions per site). See Tables 2 and 3 and the Table S1 for full strain details. Several known strains (from the NCBI database are shown in bold). The inset shows a cluster analysis based on the product type using all strains in the phylogeny.

doi:10.1371/journal.pone.0063948.g003
Table 3. Number of RFLP patterns observed per clone library.

| Library | RFLP type | n   | Closest relative in the database                  | fraction (%) |
|---------|-----------|-----|--------------------------------------------------|--------------|
| chibw1  | 1         | 44  | Weisella confuse                                  | 91.7%        |
|         | 2         | 2   | Leuconostoc pseudomesenteroides                   | 4.2%         |
|         | 3         | 1   | Weisella cibaria                                  | 2.1%         |
|         | 4         | 1   | Weisella cibaria                                  | 2.1%         |
|         | N         | 48  |                                                  |              |
| chibw2  | 1         | 30  | Lactobacillus fermentum                          | 62.5%        |
|         | 2         | 11  | Lactobacillus helveticus                         | 22.9%        |
|         | 3         | 3   | Lactobacillus harbinensis/Lactobacillus casei     | 6.3%         |
|         | 4         | 2   | Lactobacillus delbrueckii                        | 4.2%         |
|         | 5         | 1   | Acetobacter lovaniensis                          | 2.1%         |
|         | 6         | 1   | Lactobacillus fermentum                          | 2.1%         |
|         | N         | 48  |                                                  |              |
| chibw8  | 2         | 13  | Lactobacillus fermentum/Lactobacillus delbrueckii| 28.3%        |
|         | 3         | 10  | Lactobacillus delbrueckii                        | 21.7%        |
|         | 1         | 3   | Streptococcus macedonicus                        | 6.5%         |
|         | 4         | 3   | Lactobacillus fermentum                          | 6.5%         |
|         | 5         | 3   | Streptococcus macedonicus                        | 6.5%         |
|         | 7         | 3   | Lactobacillus delbrueckii                        | 6.5%         |
|         | 6         | 2   | Weisella spp                                     | 4.3%         |
|         | 8         | 2   | Lactobacillus fermentum                          | 4.3%         |
|         | 9         | 2   | Lactobacillus delbrueckii                        | 4.3%         |
|         | 10        | 2   | Lactobacillus fermentum                          | 4.3%         |
|         | 11        | 1   | Unknown                                          | 2.2%         |
|         | 12        | 1   | Weisella cibaria                                  | 2.2%         |
|         | 13        | 1   | Lactobacillus fermentum                          | 2.2%         |
|         | N         | 46  |                                                  |              |
| mab1    | 1         | 15  | Lactococcus lactis                               | 30.6%        |
|         | 2         | 13  | Streptococcus thermophiles                       | 26.5%        |
|         | 3         | 10  | Lactobacillus helveticus                         | 20.4%        |
|         | 4         | 8   | Lactobacillus helveticus; L kefirinofaciens      | 16.3%        |
|         | 5         | 2   | Lactobacillus helveticus                         | 4.1%         |
|         | 6         | 1   | Acetobacter pasteurianus                         | 2.0%         |
|         | N         | 49  |                                                  |              |
| mab10   | 1         | 26  | Lactococcus lactis                               | 55.3%        |
|         | 2         | 7   | Lactobacillus plantarum/Streptococcus equinus    | 14.9%        |
|         | 3         | 3   | Streptococcus equinus                            | 6.4%         |
|         | 4         | 3   | Streptococcus thermophiles                       | 6.4%         |
|         | 5         | 2   | Lactobacillus helveticus                         | 4.3%         |
|         | 6         | 2   | Lactobacillus brevis                             | 4.3%         |
|         | 7         | 2   | Unknown                                          | 4.3%         |
|         | 8         | 1   | Leuconostoc pseudomesenteroides                  | 2.1%         |
|         | 9         | 1   | Exiguobacterium indicum                          | 2.1%         |
|         | N         | 47  |                                                  |              |
| mab5    | 1         | 28  | Lactococcus lactis                               | 56.0%        |
|         | 2         | 10  | Citrobacter freundii/Acinetobacter ursingii      | 20.0%        |
|         | 3         | 4   | Acinetobacter spp                                | 8.0%         |
|         | 4         | 3   | Chryseobacterium bovis/Lactobacillus helveticus  | 6.0%         |
|         | 5         | 2   | Unknown                                          | 4.0%         |
|         | 6         | 2   | Lactococcus lactis                               | 4.0%         |
pooling all other clone libraries, we found a marginally significant positive relationship between phylogenetic similarity and geographical distance, just the opposite to our expectation (Figure S3).

**Discussion**

This study had two main objectives. Firstly, we characterised the microbial community structures of the products using culture based and non-culture based techniques, our primary aim being to provide and promote indigenous knowledge for potential commercial applications. The three products under investigation are widely consumed in Zambia and such basic, fundamental knowledge on the natural history of the microbial communities in these products forms a necessary first step towards understanding the details of the fermentation process and how this might be exploited to support cultural and economic development in the future. Secondly, we provide some preliminary investigations of two potential factors that shape the structure of these communities by asking whether starting raw materials or sampling location contribute to the structure of microbial communities within these products.

### 1. Microbial Community Composition

The analysis of the microbial communities in the products through culture based approaches revealed that the community in each of the product is represented by 4 to 13 dominant taxa of mainly lactic acid bacteria of genera such as *Lactobacillus, Lactococcus, Streptococcus, Weisella*, and *Leuconostoc*. This was expected since all products exhibited a pH of around 3.5 to 4.5. *Lactobacillus* and *Streptococcus* species are homofermenters producing lactic acid as major end product. *Weisella* and *Leuconostoc* species are heterofermenters that apart from acids also produce ethanol and CO₂ [3,8]. Members of the *Acetobacter* genus are the dominant gram-negative bacteria found in these products and can convert ethanol into acetic acid when enough oxygen is present [3]. The community we found is consistent with that found in other milk based products from Southern Africa [18] and other Southern and Western African cereal based products [3]. Viable counts were in the order of 10⁷ per gram of product, which is in line with viable counts found in other studies (e.g. [18]). The detection limit in our non-culture based analysis is on the order of 1 to 2% of the most abundant type since we analysed 48 RFLP patterns for each clone library.

We compared, for the same product, the microbial community structures obtained using culture and non-culture based approaches. We found that we the microbial communities among samples within the two cereal based products (Mabisi and Chibwantu) differed significantly from each other, while those for the milk based product (Mabisi) did not. While both methods give similar results at the level of the types of species found in the communities, this suggests that the laboratory culture media (solid plate count agar, MRS agar and M17 agar) quite strongly mimic the conditions in the milk products but that certain key nutrients or other conditions of the cereal products are not met. The non-culture based approach likely gives a more accurate result since this method only relies on a proper DNA extraction of all bacterial DNA from the sample and unbiased PCR [9,11]. Although it is likely that additional members may have been found for each community if we had used second generation sequencing techniques to our samples [19] the analysis of our diversity indices, rarefaction curves and estimated coverage suggests that this result is likely to be fairly robust. We conclude that there can be substantial variation among producers in the microbial community of cereal-based products despite those products having the same name and rather similar functional and organoleptic properties, although the underlying reasons why this is so remain unclear at this time.

### 2. Comparison of Microbial Community Structures between Samples Based on Differences in Raw Materials and Sampling Location

It seems reasonable to expect that the raw materials and the different fermentation procedures used to create environments with different physical properties should support and promote a different combination of microbial species. However we found only partial support for this hypothesis. Although both UniFrac significance tests and cluster analyses showed that the different products (Mabisi, Chibwantu and Munkoyo) have unique microbial communities, Munkoyo communities clustered away...
from those of both Mabisi and Chibwantu. This result is surprising because we would have expected Munkoyo and Chibwantu, being both cereal based, to be more similar to each other than either would be to Mabisi, which is milk based. Nevertheless the pH of the more similar communities, Mabisi and Chibwantu, is also very similar lending further evidence that the microbial communities are indeed quite similar. The proximate reasons for this outcome are not directly evident but may result from subtle differences in the preparation of Munkoyo and Chibwantu (Figure 1A). The boiling time for the cereal/water suspension is slightly longer for Munkoyo than for Chibwantu, for example, and the roots added are not directly evident but may result from subtle differences in the preparation at the various steps (Figure 1A, B). It may be that small differences in preparation can lead to large differences in their component microbial communities of very similar products.

We also found little support for the idea that geographical isolation among producers leads to more divergent microbial communities within a product, as would be expected to be the case if raw materials and climate are more similar for samples collected in close proximity compared to those that are more distant. The UniFrac significance tests and cluster analysis based on the populations where the samples were collected (Figure 4, Table 5) showed no signature of geographical location on community composition. This was true whether we examined all samples together or between samples of the same product (Figure 4, Table 5, Figure S3). Nevertheless there can be substantial differences in microbial community composition between producers of the same products (Tables 3 and 5), and so the question remains as to how such different communities give rise to what are recognized by consumers as effectively the same product. It may be that other factors such as soil type, microbes already present in the raw materials, or ambient climatic conditions play important roles but further tests of these hypotheses requires additional sampling and data collection.

Another, intriguing, possibility is that microbial communities of products of the same denomination are different because producers who sell their product in close proximity of each other try to achieve slightly different tastes. These different tastes, and the preferences of consumers, could be a result of the tribal background of the producer and their target consumer. This interpretation gains some support from the fact that Mumbwa, a place where we collected multiple samples, is a regional centre where different tribes congregate, although it belies the reported claims of local producers that rather uniform production procedures are employed around the country. After analysing the samples, we revisited the sampling locations to discuss the results with local producers (in Mumbwa and Lusaka). Our enquiries revealed that producers in the same town try to make their product be slightly different from others by slight modifications to the production procedure, mainly in the alteration of the incubation times at the various steps (Figure 1A, B). It may be that competition for customers among producers leads to the production of slightly different products, which would be reflected in a form of character displacement at the level of the microbial community. Given that slight differences in production methods seem to have a large effect on microbial community composition between cereal-based products, it does not seem too far-fetched to think that they might also have noticeable effects on community composition of the same product as well. Further research is required to investigate this.

The hypothesis that anthropogenic factors are important shapers of microbial communities in fermented food products is particularly exciting and could potentially explain some of the other unexpected results we obtained and deserve future exploration. Different tribes prepare the two cereal base products, Munkoyo and Chibwantu, with Munkoyo being produced primarily in the northern part of Zambia by Bemba tribes and Chibwantu being produced in central and southern Zambia by the Kaonde, Ngoni and Tonga tribes. Cultural traditions vary across Zambia [20] and variation in these traditions could become

### Table 4. Diversity indices calculated based on RFLP patterns.

| Library     | N   | n  | Chao1  | Coverage (%) | Shannon     |
|-------------|-----|----|--------|--------------|-------------|
| chibwantu1  | 48  | 4  | 4.5 (4.0–12.3)| 96           | 0.39 (0.10–0.67) |
| chibwantu2  | 48  | 6  | 6.5 (6.0–14.3)| 96           | 1.10 (0.81–1.38) |
| chibwantu8  | 46  | 13 | 13.6 (13.1–20.1)| 93          | 2.20 (1.93–2.47) |
| mabisi1     | 49  | 6  | 6*     | 98           | 1.53 (1.35–1.70) |
| mabisi10    | 47  | 9  | 9.25 (9.0–13.8)| 96          | 1.53 (1.20–1.86) |
| mabisi5     | 49  | 7  | 7*     | 98           | 1.36 (1.06–1.66) |
| munkoyo5    | 44  | 5  | 6 (5.1–18.5)| 95           | 0.77 (0.44–1.09) |
| munkoyo9    | 50  | 3  | 3*     | 98           | 0.63 (0.44–0.83) |

Numbers between brackets show 95% confidence intervals. N: number of patterns analysed; n: number of observed unique patterns; Chao1: estimation of the total number of unique types in the population; * indicates that the rarefaction curve reached saturation; coverage: fraction that our sampling covers from total number of types; Shannon: Shannon diversity index.

doi:10.1371/journal.pone.0063948.t004

### Table 5. UniFrac significance of comparisons of microbial communities based on clone library sequences in the phylogeny presented in Figure 3.

| A                      | Chibwantu | Mabisi | Munkoyo |
|------------------------|-----------|--------|---------|
| Chibwantu              | 0         | 0.03   | 0.03    |
| Mabisi                 | 0.03      | 0      | ≤0.03   |
| Munkoyo                | 0.03      | ≥0.03  | 0       |

| B                      | Chibw1    | Chibw2 | Chibw8 |
|------------------------|-----------|--------|--------|
| Chibw1                 | 0         | ≤0.03  | ≤0.03  |
| Chibw2                 | ≤0.03     | 0      | ≤0.03  |
| Chibw8                 | ≤0.03     | ≤0.03  | 0      |

| C                      | Mab1      | Mab10  | Mab5   |
|------------------------|-----------|--------|--------|
| Mab1                   | 0         | 0.99   | 0.06   |
| Mab10                  | 0.99      | 0      | 0.24   |
| Mab5                   | 0.06      | 0.24   | 0      |

| D                      | Choma     | Monze  | Mufulira | Mumbwa |
|------------------------|-----------|--------|----------|--------|
| Choma                  | 0         | ≤0.06  | 0.18     | 1      |
| Monze                  | ≤0.06     | 0      | ≤0.06    | ≤0.06  |
| Mufulira               | 0.18      | ≤0.06  | 0        | 0.06   |
| Mumbwa                 | 1         | ≤0.06  | 0.06     | 0      |

Comparison based on (A) all sequences pooled per product; (B) only the three Chibwantu libraries; (C) only the three Mabisi libraries; (D) all sequences pooled per sampling location. Values show Bonferroni corrected p-values for multiple comparisons.

doi:10.1371/journal.pone.0063948.t005
reflected in different microbial community structures of the products they produce. The role of anthropogenic and other site-specific factors in governing microbial community structure can be investigated further by including a sociological approach involving careful interviews of producers to elucidate exactly what the differences in production procedure are and investigating whether these can be linked directly to observed differences in microbial community structure.

3. Socio-economic Implications

Gaining insights into the microbiology of traditional fermented African products also has practical implications both at local and broader scales. The three fermented products that are the focus of this study are widely consumed across Zambia, however their microbial composition has to date not been studied. Knowledge on the microbial composition and factors that affect this composition will promote and improve indigenous knowledge and practices, enhance food security for local communities at low costs [21] and provide opportunity for economic development.

Fermented products are known to promote human health because they are nutritious and safe: the microbial communities in the products are highly stable and are able to withstand an invasion by pathogenic bacteria and in some cases detoxify the raw materials [8,21,22]. We found that the microbial communities in the products are predominated by lactic acid bacteria, which supports the claim by local knowledge that consumption of Mabisi, Chibwantu and Munkoyo promotes human health. In particular it is claimed that these products suppress diarrhea and are thought to mitigate symptoms of HIV/AIDS and hypertension and anti-allergen properties [23], although more direct tests of these claims is warranted. Nevertheless, lactic acid bacteria are known for their specific antimicrobial properties that determine their bio-preservation properties in foods, including the production of organic acids, hydrogen peroxide, carbon dioxide, diacetyl and broad-spectrum antimicrobials such as anti-competitor toxins [8].

Traditional products, including how exactly their safety is ensured, could further point to ways to optimize existing fermentation processes, to novel processes which may provide new strategies to fight more serious infections of worldwide concern, and facilitate the effective marketing the traditional version of the products, which can generate income for local communities [1,25].

4. Traditional Fermented Products as Model for the Study of Ecosystem Ecology and Evolution

Microbial systems are now routinely used to study the basic evolutionary processes of adaptation and diversification in well-defined communities of one or at most a few species [4,26,27,28,29,30]. Ideally, scientists studying evolution would like to predict the ecological properties and evolutionary dynamics of highly complex ecosystems as well but the ability to scale up their inferences from simple communities in the lab to more complex ones in nature remains limited. Needed to fill this gap are experimental model systems for studying long-term whole community evolution, but these are almost entirely lacking [31,32]. The products used in this study may be such a system. Since the product is produced by repeated propagation of starter culture and without exchange with other producers, the constituent microbial communities have been evolving independently from each other probably for at least hundreds, if not thousands, of microbial generations. In essence, then, each product from a different producer represents an independent instance of adaptive community evolution towards a defined end-point – the fermented product – with known ecological function. These functions include both nutritional content, appeal to the consumer, and safety. This system thus opens new avenues for research on how complex communities with known functional properties are assembled and evolve.

Supporting Information

Figure S1 Rarefaction curves of clone libraries based on RFLP patterns.

Figure S2 RFLP patterns of 48 clones from the Mabisi clone library, digested with EcoRI and HaeIII. The ladder is a 100 bp ladder. From these patterns, the number and frequency of the different distinct patterns observed here was used to calculate diversity indices. In this library, we observed 6 different patterns. To the right of each row is the non-digested negative control.

Figure S3 Phylogenetic similarity of microbial communities as a function of geographical distance. We
examined the relationship between the geographical distance at which the samples were collected and the similarity of the microbial community composition; this similarity was computed using the phylogenetic tree using one library (Chibwantu1) as a reference. Pooling all clone libraries, we found a marginally significant positive relationship between phylogenetic similarity and geographical distance ($r^2 = 0.5385$, $F_{1,5} = 5.83$, $p = 0.060$). Since one library is used as a scaling reference in each comparison, the analysis for each product type could only be done with the Chibwantu and Mabisi libraries, for which we each analysed three samples, leaving us with only two points for each library and thus no statistical power to detect a relationship between phylogenetic similarity and geographical distance for each product type separately. Nevertheless, this analysis confirms the analysis described above which showed that phylogenetic similarity is lowest for microbial communities in samples collected in close proximity. A generalised linear model on the pooled dataset with phylogenetic similarity as the dependent variable and geographic distance and product type as independent variables revealed no significant effect of either independent variable nor of their interaction on the variation in phylogenetic similarity. (TIF)

Table S1 Isolates and clones most closely related relatives in NCBI and RDP databases. (XLSX)

Acknowledgments

We thank Thera Rasing, Shannon Watson, Maggie Stoeye, Christina Hoffing and John Shindano and two anonymous Reviewers for help, comments and assistance.

Author Contributions

Conceived and designed the experiments: SS CK RK AJP. Performed the experiments: SS CK CT. Analyzed the data: SS AJP. Contributed reagents/materials/analysis tools: RK AJP. Wrote the paper: SS CK RK AJP.

References

1. Odumé SA (1980) African fermented foods: from art to science. MIRCEN journal of applied microbiology and biotechnology 4: 259–273.
2. Nout MJR, Darkar PK, Beuchat LR (2007) Indigenous fermented foods. In: Doyle MP, Beuchat LR, editors. Food Microbiology: Fundamentals and Frontiers, 3rd edition. Washington DC: ASM Press, pp.817–835.
3. Blandino A, Al-Aseeri ME, Pandiella SS, Cantero D, Webb C (2003) Cereal-based fermented foods and beverages. Food Research International 36: 527–543.
4. Lawrence D, Figuia F, Belrendes V, Bundy JG, Phillimore AB, et al. (2012) Species Interactions Alter Evolutionary Responses to a Novel Environment. Plos Biology 10.
5. Nguz K, Kasase C, Phiri G, Shindano J, Khizezi DM (2004) Isolation and inhibitory effects of lactic acid bacteria from selected traditional fermented beverages from rural Zambia on some food pathogens. Journal of Science and Technology (Zambia) 1: 9–16.
6. Zulu RM, Dillon VM, Owens JD (1997) Munkoyo beverage, a traditional Zambian fermented maize gruel using Rhynchosia root as amylase source. International Journal of Food Microbiology 34: 249–258.
7. Haard NF, Oduma SA, Quintero-Ramirez R, Lorenz-Quinones A, Wacher-Radaire C (1999) Fermented Cereals: a global perspective. Food and Agriculture Organisation.
8. Caplice E, Fitzgerald GF (1999) Food fermentations: role of microorganisms in food production and preservation. International Journal of Food Microbiology 50: 131–149.
9. Abriouel H, Ben Omar N, Lopez RL, Martinez-Camamero M, Keleke S, et al. (2006) Culture-independent analysis of the microbial composition of the African traditional fermented foods poto poto and degue by using three different DNA extraction methods. International Journal of Food Microbiology 111: 228–233.
10. Ampe F, Ben Omar N, Moizan C, Wacher C, Guyot JP (1999) Polyphasic study of the spatial distribution of microorganisms in Mexican pozol, a fermented maize dough, demonstrates the need for cultivation-independent methods to investigate traditional fermentations. Applied and Environmental Microbiology 65: 5464–5473.
11. Ercolini D, Moscheti G, Blaostia G, Coppola S (2001) The potential of a polyphasic PCR-DGGE approach in evaluating microbial diversity of natural whey cultures for water-buffalo mozzarella cheese production: Bias of culture-dependent and culture-independent analyses. Systematic and Applied Microbiology 24: 610–617.
12. Cole JR, Wang Q, Cardenas E, Fish J, Chai B, et al. (2009) The Ribosomal Database Project: improved alignments and new tools for rRNA analysis. Nucleic Acids Research 37: D141–D145.
13. Schloss PD, Westcott SL, Ryabin T, Hall JR, Hartmann M, et al. (2009) Introducing mothur: Open-Source, Platform-Independent, Community-Supported Software for Describing and Comparing Microbial Communities. Applied and Environmental Microbiology 75: 5421–5425.
14. Ashelford KE, Chuzhanova NA, Fry JC, Jones AJ, Weightman AJ (2006) New screening software shows that most recent large 16S rRNA gene clone libraries contain chimeras. Applied and Environmental Microbiology 72: 5734–5741.
15. Posauda D (2008) jModelTest: Phylogenetic model averaging. Molecular Biology and Evolution 25: 1253–1266.
16. Price MN, Dehal PS, Arkin AP (2009) FastTree: Computing Large Minimum Evolution Trees with Profiles instead of a Distance Matrix. Molecular Biology and Evolution 26: 1641–1650.
17. Josopone C, Hamardy M, Knight R (2006) UniFrac - An online tool for comparing microbial community diversity in a phylogenetic context. BMC Bioinformatics 7.
18. Beukes EM, Bester BH, Mostert JF (2001) The microbiology of South African traditional fermented milks. International Journal of Food Microbiology 65: 189–197.
19. Pogacic T, Kelava N, Zamberlin S, Dolenic-Spehar I, Zambrsija D (2010) Methods for Culture-Independent Identification of Lactic Acid Bacteria in Dairy Products. Food Technology and Biotechnology 48: 3–10.
20. Sojkowski J (2006) Culture of Zambia. September 2006 ed.
21. Mensah P (1997) Fermentation - the key to food safety assurance in Africa? Food Control 8: 271–278.
22. Odumé AO, Oyewole OB, Oduamau AO (2009) Microbiological safety and quality assessment of some fermented cassava products (afihan, fufu, garri). Journal of Food Science and Technology 4: 432–435.
23. El-Ghaish S, Ahmadova A, Hadji-Stani I, El Mecherfi K, Bazaniky I, et al. (2011) Potential use of lactic acid bacteria for reduction of allergenicity and for longer conservation of fermented foods. Trends in Food Science & Technology 22: 509–516.
24. Adam MR, Moss MO (2004) Food Microbiology, 2nd edition. Cambridge, UK: The Royal Society of Chemistry.
25. Oyewole OB (1997) Lactic fermented foods in Africa and their benefits. Food Control 8: 289–297.
26. Kassen R, Rainey PB (2004) The ecology and genetics of microbial diversity. Annual Review of Microbiology 58: 207–231.
27. Chao I, Levin BR, Stewart FM (1977) Complex community in a simple habitat - experimental study with bacteria and phage. Ecology 58: 369–378.
28. Jacks JP, Forde SE, Bohman BM (2005) Microbial experimental systems in ecology. Advances in Ecological Research, Vol 37: Population Dynamics and Laboratory Ecology 37: 273–307.
29. Meyer JR, Schoustra SE, Lachapelle J, Kassen R (2011) Overshooting dynamics in a model adaptive radiation. Proceedings of the Royal Society B-Biological Sciences 278: 392–398.
30. Rainey PB, Travisano M (1998) Adaptive radiation in a heterogeneous environment. Nature 394: 69–72.
31. Curtis TP, Sloan WT (2004) Prokaryotic diversity and in limicorl microbial community structure in nature and implications for microbial ecology. Current Opinion in Microbiology 7: 221–226.
32. Jessop CM, Kassen R, Forde SE, Kerr B, Buckling A, et al. (2004) Big questions, small worlds: microbial model systems in ecology. Trends in Ecology & Evolution 19: 189-197.