Introduction

The oral cavity is a complex human habitat which is home to more than 750 different bacterial species [1]. Many of these bacteria are considered as members of the physiological flora living in symbiotic or commensal relationship to each other and their host. However, several bacterial species are pathogenic and have been reported as causative agents for oral diseases like caries, gingivitis, and periodontitis [1].

In most niches of the oral cavity physiological and pathogenic species live in large highly structured communities called biofilms [2–4]. Such biofilms develop after adherence of primary colonizing bacteria to inert or chemically conditioned surfaces. During biofilm maturation new bacterial species bind as secondary colonizers on top of the pioneer layers. Structure and integrity of biofilms is secured by production of macromolecular extracellular substances. Depending on the species composition an intense, inter- and intra-species communication occurs in such oral biofilms [5,6]. Oral microbial biofilms may contain several hundred species and their composition can significantly differ among individuals [7,8].

It is now appreciated that biofilm-involving oral diseases are among the most prevalent diseases worldwide, affecting nearly every human being at some time point during life span [9,10]. Thus, investigation and comprehension of initiation and progression of these diseases, as well as developing novel strategies and therapies to control such biofilm-associated oral diseases is of great scientific interest and importance.

Numerous in vitro and in vivo studies have elucidated physiological and molecular details of bacterial interactions in oral biofilms of pathogenic and commensal bacteria. Such studies identified critical virulence factors of single species and allowed a first insight on the complex pathomechanisms acting in diseases like caries, gingivitis, and periodontitis [11–16]. It is now clear that mechanisms like cross-feeding and metabolic cooperation, as well as intra-/inter-species communication via quorum-sensing are critical for oral disease initiation and progression [3].
One level of influence in poly-microbial interactions is gene transcription. RT-qPCR and DNA-array design, based on full genome sequences, have boosted studies on expression profiles of oral bacteria. Specifically, single-species investigations employing typical stress forms encountered by the bacteria in their natural habitat were performed [17–19], planktonic versus biofilm bacterial growth was compared, and wild type and mutant strains were analyzed in such comparative transcriptomic approaches [20–22].

However, as the consortium of bacteria in the oral cavity is a multi-species association it would be an advantage and progress to analyze different oral bacteria with one array. Since to the best of our knowledge such a multi-species array was not previously available for transcriptome studies of oral bacteria, we developed a five-species custom array based on NimbleGen® DNA array platform technology. The 385 K™ chip design, allowing a total number of 385,000 gene probes per array, appeared to be a well suited platform to cover several bacterial genomes with up to three replicates per probe on each array. Our novel array design covered the full genomes of the cariogenic species Streptococcus mutans UA159, the physiological species Streptococcus sanguinis SK36, and the three periodontitis-associated species Fusobacterium nucleatum DSMZ 25586, Porphyromonas gingivalis W83, and Aggregatibacter actinomycetemcomitans HK1651. The chip contained 10,107 unique probe sets covering 10,186 genes. Most genes were covered by 13 individual probes.

Here we present transcriptome data of single-species cultures and biofilms, which allowed identification of cross-hybridizing single gene probes across the species represented on the array. As an example, the transcriptome of S. mutans under the influence of S. mits was studied in a two-species biofilm. Array results were verified by real-time RT-qPCR.

This array is a powerful tool to investigate and understand the complexity of transcriptional changes in mixed-species biofilms and could be useful to study biofilm interfering substances and their mode of action.

**Results**

Single-species hybridizations on the five-species arrays allowed identification of cross-hybridizing gene probes

In advance of using this novel custom array design for multi-species experiments, it was mandatory to perform hybridizations with labeled cDNAs from single-species cultures and biofilms. Under the experimental setup described here, only S. sanguinis and S. mutans developed biofilm structures. All other tested species grew in the planktonic phase or dense monolayers in the culture flasks. A documentation of biofilm and monolayer formation of the single species is given in Supporting Figure S1.

Hybridization of labeled cDNA from a single species resulted in fluorescence signals which can be presented in three groups: i) the total number of fluorescence signals compared to total number of probes derived from the whole genome, ii) probes reacting with different fluorescence signal intensities, and iii) nonspecifically reacting probes derived from the other bacterial species.

The hybridization of labeled cDNA from 24 h S. sanguinis biofilms resulted in fluorescence signals for 9497 probes, representing 1550 transcriptionally active genes. Thus, 69.1% of the total number of 2244 genes is transcribed under the biofilm growth conditions used in this study. One gene was classified as highly transcribed (6 probes of this gene reached 75–100% of maximal fluorescence signal intensity of the array). Up to now this gene has been described to encode a conserved hypothetical protein.

Further, among 38 probes belonging to 14 genes with fluorescence intensities of 50–75% above background signal, we found many chaperones: dnaK [SSA_2007], dnaJ [SSA_2005], grpE [SSA_2008], clpP [SSA_2096], responsible for protein folding and degradation, ftsH [SSA_0015], a Zn metallo-peptidase involved in cell division via sigma 32, adcC [SSA_0136] a putative Zn ABC transporter, and tptN [SSA_0117], a ribosomal protein (Supporting Table S1).

Besides, 58 probes were identified to give false positive signals. These probes were derived from 9 genes of A. actinomycetemcomitans HK1651, 5 genes of S. mutans UA159, 1 gene of P. gingivalis W83 and 1 gene of F. nucleatum ATCC 25586. Table 1 summarizes the numbers of probes and designations of corresponding genes, which were found to lead to nonspecific signals after single species hybridization of all bacterial species presented on the chip. More detailed information on these probes and genes is presented in Supporting Table S2. Figure S1a shows the signal intensities graphically. However, since most genes were covered by up to 13 probes on the array there is only in rare cases a need to fully exclude complete genes from the analysis. For example, S. sanguinis cDNA cross-hybridized with the gene probe AA00182P0047 from A. actinomycetemcomitans HK1651 (see Supporting Table S2).

| Hybridization with: | S. mutans | S. sanguinis | P. gingivalis | F. nucleatum | A. actinom. |
|---------------------|-----------|--------------|--------------|--------------|------------|
| S. mutans (1927 genes) | 24210 probes | 187 probes | 16 probes | 11 probes | 68 probes |
| S. sanguinis (1915 genes) | 14 probes | 9497 probes | 1 probe | 1 probe | 42 probes |
| P. gingivalis (2244 genes) | (5 genes) | (1550 genes) | (16 genes) | (11 genes) | (17 genes) |
| F. nucleatum (1883 genes) | 8 probes | 12 probes | (1771 genes) | (4 genes) | (13 genes) |
| A. actinom. (1964 genes) | 5 probes | 22 probes | 3 probes | 5223 probes | 24 probes |
| A. actinom. (2168 genes) | 9 probes | 12 probes | 8 probes | 7 probes | 6702 probes |

Table 1. The number of cross-hybridizing probes.

Identification of non-specific hybridizing probes after single hybridization of S. mutans UA159, S. sanguinis SK36, P. gingivalis W83, F. nucleatum ATCC25586 and A. actinomycetemcomitans HK1651.

The table shows cross-hybridizing probes affecting different numbers of genes (in brackets) after single-species hybridization. Results which refer to the species hybridized on the chip are displayed in bold characters.

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Supporting Table S3 revealed that the gene AA00182 is covered by 13 individual gene probes. Thus, 12 non-crossreactive gene probes covering AA00182 are still available for evaluation of differential expression in an A. actinomycetemcomitans HK1651 - S. sanguinis mixed-species setting (Table 2).

Chip-hybridization with cDNA from S. mutans UA159 24 h biofilms revealed 99.4% of the genome actively transcribed under the experimental conditions described here. 24,210 probes of S. mutans UA159, representing 1915 genes from a total number of 1927 genes, were detected. Among the highly abundant transcripts (436 probes covering 167 genes with 75–100% of the maximal fluorescence signal intensity of the array), we identified also chaperones like dnaK (SMU_82), dnaJ (SMU_83), and groEL (SMU_1594). Further, ftsZ (SMU_552), a putative cell division protein, dnaJ (SMU_1921), coding for a DNA helicase, pflA (SMU_1692), coding for pyruvate formiat lyase A, and several genes encoding ribosomal proteins like S7 (SMU_358), L27 (SMU_849), S1 (SMU_1200), S11 (SMU_2002) were found (Supporting Table S1). A total of 111 genes from the other species on the chip were found to react nonspecifically (refer to Table 1, Supporting Table S2 and Supporting Figure S2a). Again, with the information from Supporting Table S2 (number and identity of individual cross-hybridizing probes per gene) and the information from Supporting Table S3 (total number of individual probes per gene on the array) the experimentalist can decide if whole genes need to be excluded from two-species analysis.

The hybridization with labeled cDNA of P. gingivalis W83 cultures revealed measurable fluorescence intensities for 21855 genes, representing 1165 of 2138 genes of this species. Particularly, 8 genes, represented by 54 probes, were found to be highly transcribed (75–100% of the maximal fluorescence signal intensity of the array). These genes encoded chaperones like dnaK (PG_1208), groES (PG_0521), groEL (PG_0521), clpP (PG_0010), clpX (PG_0417) and several ribosomal proteins, like L27 (PG_0315), S9 (PG_376), L1 (PG_391), and S6 (PG_595) (Supporting Table S1). Only 75 probes originating from 24 genes of the other species reacted nonspecifically (refer to Table 1, Supporting Table S2 and Supporting Figure S2b).

For F. nucleatum DSMZ 25586, which grew in dense monolayers after 24 h incubation, 25506 fluorescence signals were obtained for 5223 probes belonging to 1614 of 1646 genes of this species (92.2% of the genome). Only 7 genes, represented by 7 probes, showed very high fluorescence intensities, suggesting only a small number of genes as highly transcribed under the chosen conditions (Supporting Table S1). Among them, we identified genes encoding for a malonyl-CoA transacylase (FN0149), acetyl-CoA acetyltransferase (FN0495), a alkyl hydroperoxide reductase (FN1983) and also ftsZ (FN1451) involved in cell division. Further, 12 genes of A. actinomycetemcomitans HK1651, 15 genes of S. sanguinis SK36, 4 genes of S. mutans UA159 and 5 genes of P. gingivalis W83 displayed nonspecific reactivities (refer to Table 1, Supporting Table S2 and Supporting Figure S2d).

Hybridization of A. actinomycetemcomitans labeled cDNA resulted in fluorescence signals for 6702 probes of A. actinomycetemcomitans, representing 1165 of 2138 genes of this species. Particularly, 8 genes, represented by 54 probes, were found to be highly transcribed (75–100% of the maximal fluorescence signal intensity of the array). Except for comB (AA00059), a gene encoding for a home exporter protein, all other genes have unknown or hypothetical functions (Supporting Table S1). 36 probes led to nonspecific signals potentially affecting the analysis of 15 genes of the other species presented on the chip (refer to Table 1, Supporting Table S2 and Supporting Figure S2e).

Taking into account the relatively low number of nonspecific cross-species hybridization signals identified in this series of experiments, the initial sequence analysis and algorithms for probe design proved to be of high quality. In only 3 cases, all probes derived from one gene showed nonspecific signals and thus, would require the exclusion of these genes from further analysis in the respective two-species setting (listed in Table 2). After performing these corrections, the combination of cDNAs from two or more species of the array should lead to truly positive signals in all possible combinations.

**Table 2. Exemplary genes including cross-hybridizing probes.**

| Gene ID | Probe ID |
|---------|----------|
| AA00182 | AA00182P00018 AA00182P00311 AA00182P00470 | AA00182P00623 AA00182P00654 AA00182P00739 |
| AA00182 | AA00182P00018 AA00182P00311 AA00182P00470 | AA00182P00623 AA00182P00654 AA00182P00739 |
| AA00182 | AA00182P00018 AA00182P00311 AA00182P00470 | AA00182P00623 AA00182P00654 AA00182P00739 |
| SSA_0123 | SSA_0123P00062 SSA_0123P00101 SSA_0123P00105 | SSA_0123P00110 SSA_0123P00115 SSA_0123P00120 |
| SSA_0440 | SSA_0440P00032 SSA_0440P00039 SSA_0440P00042 | SSA_0440P00046 SSA_0440P00049 SSA_0440P00052 |
| SSA_2391 | SSA_2391P00012 SSA_2391P00013 SSA_2391P00014 | SSA_2391P00015 SSA_2391P00016 SSA_2391P00017 |

| Probe ID | AA00182P000851 AA00182P001106 AA00182P001117 AA00182P001129 AA00182P001135 AA00182P001141 |
|----------|-------------------------------------------------|
| SSA_0123P00222 | SSA_0123P00238 SSA_0123P00237 SSA_0123P00238 SSA_0123P00240 SSA_0123P00243 SSA_0123P00296 |
| SSA_0440P00053 | SSA_0440P000174 SSA_0440P000177 SSA_0440P000178 SSA_0440P000179 SSA_0440P00180 SSA_0440P00181 |

Exemplary genes and their probes represented on the described array. Cross-hybridizing probes are indicated by bold characters. Footnotes indicate probes which were positive after hybridization with cDNA from P.gingivalis (†), S. mutans (‡) and S. sanguinis (§), respectively. doi:10.1371/journal.pone.0027827.t002
study from our lab has documented that the cariogenic species *S. mutans* grows to significantly increased biofilm mass in a dual culture with the physiological species *S. mitis* over a five day investigation period [23]. At the time of array design no full *S. mitis* genome sequence was available, thus this species is not covered by the array. However, the taxonomic relationship between both species could interfere with the suitability of any array to analyze the specific transcriptome of the two species. Thus, with the next series of experiments we addressed several issues: i) was the design of the present array suitable to discriminate the transcriptomes of two closely related species, ii) could this issue be tackled in absence of the genome sequence of one species, and iii) would the array hybridization experiments render results which could explain the observed behavior of *S. mutans* mixed-species biofilms? Therefore, we first tested potential cross-reactions of *S. mitis* cDNA with the 5 species genome array by exclusively using *S. mitis* cDNA. In this assay, 311 *S. mutans* probes displayed nonspecific reactions (Supporting Table S4). These nonspecifically reacting probes, but not the whole genes, were excluded from the data analysis of the *S. mutans/S. mitis* single-species/co-culture experiments.

Under the chosen conditions, *S. mutans* was able to form large biofilm masses already after 24 hours of incubation. The combination of *S. mutans* with *S. mitis* resulted in larger biofilm masses than obtained for the *S. mutans* mono-species culture. Culture supernatant pH-values of the combination were lower than for the *S. mutans* mono-species culture (4.96 versus 5.8 after 24 hours) but numbers of colony forming units of *S. mutans* were unchanged within the boundaries of experimental error over a five day co-culture period with *S. mitis* [23]. In contrast, *S. mitis* could be detected via different microscopic techniques, but no colony forming units were found.

After deletion of all nonspecifically reacting probes from further analysis, the *S. mutans* transcriptome after co-culture with *S. mitis* showed a significantly different transcript abundance for 30 genes (fold change two, p<0.05) (19 were found up-regulated, 11 were down-regulated) compared to the *S. mutans* mono-species biofilm transcriptome. Using the Internet platform KEGG (Kyoto Encyclopedia of Genes and Genomes, http://www.genome.jp/kegg/), we found that the majority of these differentially regulated genes were involved in metabolism and environmental information processing (membrane transport) (Table 3). Of note, the stress response operon *dnaK* (smu_80-83) and the *cop*-operon (SMU_424-smu_427), as well as *groEL* (SMU_154) and *clpL* (SMU_956), which have been described to be involved in stress response, showed high transcript abundances (Figure 1). Moreover, the agmatine deiminase gene cluster [24] (SMU_262-264) was found downregulated.

The microarray data have been deposited in the Gene Expression Omnibus (GEO) database under accession number GSE28041. 

**Confirmation of transcriptome results by Real Time RT-qPCR**

As an independent second method we used RT-qPCR for validation of the observed transcriptome data. A variety of genes showing high or low transcript abundances in transcriptome analysis were chosen for further analysis with cDNA from a new series of biologically independent experiments. Precisely, 7 up-regulated and 6 down-regulated genes were selected. The gene SMU_2013 was used as control as it showed no change of expression (p>0.95) in the array experiments.

Table 4 shows the results of RT-qPCR as well as the comparison to the data of transcriptome analysis. In general, RT-qPCR confirmed the results of transcriptome experiments.

**Discussion**

Oral microbial diseases are based on qualitative and quantitative dysbalances of the physiological and pathogenic microbial flora and thus are polymicrobial by nature. Comparability of current *in vitro* investigations on microbial gene expression during health and diseased status is limited due to usage of diverse experimental setups. Differences include environmental conditions as well as static or flow through culture systems. The informative value of bacterial transcriptome studies is restricted in most cases due to the single-species DNA array setup. To overcome some of these shortcomings we here describe the setup and use of a 5-species transcriptome array for the study of both single- and multi-species cultures and biofilms. Only a minority between 0.35% (in case of *P. gingivalis*) and up to 1.15% (in case of *S. mutans*) of probes displayed unspecific hybridization results. Due to the high redundancy of probes per gene, these unspecific probes could be excluded from further analysis while the gene could still be included in the analysis in all but 3 cases.

In a first series of experiments, we simultaneously identified unspecific probes on the array and measured gene expression of early single-species planktonic cultures, monolayers and biofilms. Corresponding to the status of young biofilms, transcriptome analysis from the contained cells demonstrated the presence of transcripts of the vast majority of genes, especially such encoding all chaperones and ribosomal proteins, for example *dnaK* (SSA_2007; SMU_0082; PG_1208; AA000657) in *S. sanguinis*, *S. mutans*, *P. gignicilis* or *A. actinomycetemcomitans*, *clpL* (SSA_2096; PG_0010) in *S. sanguinis* and *P. gingivalis*, or a member of the *fts* operon, involved in cell division (SSA_0015; SMU_552; FN1451) in *S. sanguinis*, *S. mutans* and *F. nucleatum*. The ubiquitous presence of chaperone transcripts could be an indicator for general or acidic stress in these cultures [17,25–27].

A complete *S. mitis* genome sequence was not available at the time of the array design, therefore the corresponding probes were not included. However, it was of special interest whether the array could still be used for a two-species analysis with a species represented on the chip and another that was not included. Again, specificity testing with *S. mitis* cDNA demonstrated cross hybridization of only a few *S. mutans* probes, allowing the inclusion of all *S. mutans* genes in the final analysis. The two-species combination was phenotypically described previously under growth conditions which supported planktonic growth of both species, but biofilm formation only for *S. mutans* [23]. The glucose-free growth medium CDM/sucrose is a prerequisite to mimic cariogenic biofilms with thick extracellular polysaccharides [28]. Under such virulent cariogenic conditions previous studies have shown suppression of *S. sanguinis* growth in co-culture with *S. mutans* [29].

Apparently, *S. mutans* suppressed *S. mitis* culturability in the biofilm, however, initial presence of *S. mitis* as a physiological species increased the cariogenic potential of the *S. mutans* biofilm. Does differential gene expression explain any of these phenotypes? For example, the co-cultivation of *S. mitis* with *S. mutans* resulted in high transcript abundances of *gfpE* (SMU_81). However, only the inspection of the whole operon (SMU_80-83) allowed the determination that all genes of this operon were found in higher transcript abundances compared to the *S. mutans* mono-species culture. In general, results of transcriptome analysis of the *S. mitis*/ *S. mutans* co-culture were in good agreement with the former obtained functional results [23]. Stan dar et al. showed that the co-cultivation of both species led to higher biofilm masses compared to the *S. mutans* mono-species culture, whereas pH was lower in the *S. mitis*/ *S. mutans* combination. Transcriptome analysis revealed
that several genes encoding stress proteins of *S. mutans* were upregulated under the co-cultivation conditions. Earlier studies indicated an association between expression of *S. mutans* stress proteins at a lower pH-value. For instance, acid stress/shock induced transcription of the molecular chaperons *dnaK* and *groEL* [25,27], which agreed with our findings in transcriptome and RT-qPCR analysis. Both proteins play essential roles in cellular metabolism by assisting in the folding of newly synthesized or denatured proteins, as well as in the assembly, transport, and degradation of cellular proteins [30]. Furthermore, HrcA, the repressor of class I heat shock gene expression and encoded by the first gene of the *dnaK* operon of *S. mutans* (*hrcA-grpE-dnaK*) negatively regulates the *dnaK* and the *groE* operon [27]. In the present study the *hrcA* gene transcript was decreased (*p* = 0.058), consistent with the increased presence of the *groE* and *dnaK* operon transcripts.

Also the *clp* – system was shown to play an important role in acid tolerance [17,26]. The genome of *S. mutans* UA159 encodes
orthologs of five Clp ATPases (ClpB, ClpC, ClpE, ClpL, and ClpX), and a single ClpP peptidase [17,31]. A clpL knock-out mutant of S. mutans displayed a raised sensitivity against acid killing [26]. Other studies could demonstrate that a ClpL homologue interacts with the HSP70 system, which includes DnaK and DnaJ to form a chaperone complex [32]. Therefore, one could speculate that ClpL plays an important role in acid stress tolerance. Here we could measure increased clpL transcript abundance in the presence of a low pH in the S. mitis/S. mutans co-cultures. In good agreement with previous studies on S. mutans acid stress behavior [33], our study of the S. mitis/S. mutans co-culture revealed the upregulation of copY and copZ. In addition, the copYAZ

![Figure 1. Overview of transcript level changes of S. mutans in co-culture with S. mitis compared to its single-species culture.](image)

Table 4. Verification of array results by RT-qPCR analysis of selected genes.

| Gene ID  | Gene name/Description                                      | Fold change of RT-PCRa | Fold change of transcriptome analysisb |
|---------|-----------------------------------------------------------|-------------------------|----------------------------------------|
| SMU_81  | grpE; heat shock protein GrpE (HSP-70 cofactor)           | 2.04 (0.51)             | 2.96 (p<0.05)                          |
| SMU_312 | PTS system, sorbitol phosphotransferase enzyme IIIBC      | 6.12 (1.25)             | 2.92 (p<0.05)                          |
| SMU_426 | copA; copper-transporting ATPase; P-type ATPase           | 3.25 (0.83)             | 3.31 (p = 0.072)                       |
| SMU_956 | clpL; putative Clp-like ATP-dependent protease, ATP-binding subunit | 5.11 (3.3)             | 3.87 (p = 0.069)                       |
| SMU_1495| lacB; galactose-6-phosphate isomerase, subunit LacB      | 2.09 (0.52)             | 2.03 (p = 0.15)                        |
| SMU_1954| groEL; putative chaperonin GroEL                          | 2.15 (0.83)             | 2.33 (p = 0.075)                       |
| SMU_2047| ptsG; putative PTS system, glucose-specific IIABC component | 3.88 (2.79)             | 2.13 (p<0.05)                          |

upregulated genes

| Gene ID  | Gene name/Description                                      | Fold change of RT-PCRa | Fold change of transcriptome analysisb |
|---------|-----------------------------------------------------------|-------------------------|----------------------------------------|
| SMU_262 | otcA; putative ornithine carbamoyltransferase             | –2.39 (1.55)            | –2.07 (p<0.05)                         |
| SMU_498 | conf; putative late competence protein                   | –2.03 (0.93)            | –2.16 (p<0.05)                         |
| SMU_772 | gldP; glucan-binding protein D with lipase activity; BglB-like protein | –3.06 (0.06)             | –2.8 (p<0.05)                         |
| SMU_1055| radC; putative DNA repair protein RadC                   | –1.64 (0.45)            | –2.35 (p<0.05)                         |
| SMU_1340| bacA2; putative surfactin synthetase                     | –3.36 (0.39)            | –1.87 (p = 0.054)                      |
| SMU_1934c| putative cobalt ABC transporter, ATP-binding protein    | –1.99 (0.48)            | –2.07 (p<0.05)                         |

downregulated genes

Data are from three biological replicates.

aAverage n-fold change in expression for three biological and three technical replicates, each. Values in parentheses are standard deviations.
bn-fold change after expression array analysis of three biological replicates. Significance is indicated by p-values in parentheses.

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A Five-Oral Species DNA Array

In summary, we have developed a 5-species transcriptome array based on NimbleGen™ 305 K array technology. We showed that the chosen array setup can reliably detect the transcript abundance of whole genomes in single-species experiments. The obtained results uncovered genes which are actively transcribed under the defined environmental and culture conditions. Additionally, none specifically hybridizing probes were identified. They can be excluded from further analysis in multi-species investigations based on the individual specificity parameters and requirements of the researcher, however, in most cases without losing the whole gene for the analysis. To our best knowledge, so far there is no array available for multi-species transcriptome analysis. Therefore, the newly designed chip could be a powerful tool to simultaneously investigate the transcriptome of more than one species from the oral environment.

Materials and Methods

Bacterial strains and culture conditions

The bacterial strains Streptococcus mitis ATCC 11843, Streptococcus mutans UA159, S. sanguinis SK36, F. nucleatum ATCC 25586, P. gingivalis W83 and A. actinomycetemcomitans HK1651 were purchased from commercial providers (DSMZ, Braunschweig, Germany and ATCC, Manassas, USA). Streptococcus mitis ATCC 11843, Streptococcus mutans UA159, S. sanguinis SK36 and A. actinomycetemcomitans HK1651 were cultured in brain heart infusion medium (BHI; Oxoid) at 37°C under a 5% CO₂-20% O₂ atmosphere. F. nucleatum ATCC 25586 and P. gingivalis W83 were cultivated in BHI supplemented with 0.25% glutamate or 5 µg/ml hemin and 50 mM galactose, respectively, at 37°C under a 10% CO₂ – 10% H₂ – 80% N₂ atmosphere.

Setup of adherent cell cultures and biofilms

Bacteria were grown in BHI to stationary phase, washed with phosphate buffered saline (PBS, pH 7.4), and adjusted to a strain specific OD₆0₀ to obtain 1×10⁸ cells ml⁻¹. Subsequently, each bacterial suspension was diluted 10-fold in a chemical defined medium (CDM) supplemented with 50 mM sucrose (S. mutans, S. sanguinis, A. actinomycetemcomitans) or in artificial saliva supplemented with 50 mM galactose (P. gingivalis and F. nucleatum) and inoculated in 75 cm² CELLSTAR® tissue culture flasks (Greiner bio-one, Frickenhausen, Germany). The bacteria were cultured alone to establish species-dependent mono-species cultures and biofilms or in combination resulting in two-species biofilms. All cultures were grown in an anaerobic incubator under an appropriate atmosphere (80% N₂, 10% CO₂, 10% H₂) at 37°C for one day under static conditions. The incubator atmosphere was saturated with water vapor to prevent exsiccation of the cultures and it was constantly exposed to a platinum catalyst to decrease the content of short-chained fatty acids in the atmosphere. Monolayer formation and biofilm growth was determined as described previously [23].

RNA preparation from whole biofilm samples, cDNA synthesis, labeling and hybridization

To obtain samples for RNA preparation, mono- and two-species cultures and biofilms were set up in uncoated CELLSTAR® issue culture flasks (75 cm², 250 ml) (Greiner Bio-One, Frickenhausen, Germany) and allowed to build for one day of incubation under anaerobic conditions. In the cases when an isolate only formed a cell monolayer, the culture supernatant and the cells removed from the monolayers were combined to achieve a significant cell mass for RNA preparation (A. actinomycetemcomitans, F. nucleatum, P. gingivalis). For species with the capacity to
form thick biofilms (S. sanguinis, S. mutans) under the chosen assay conditions, liquid medium was removed and biofilms were gently washed with PBS. Biofilms were detached from the surface by thorough scraping and washing with PBS. The biofilm suspension was centrifuged and the obtained pellet was used immediately or frozen at −80°C until further use.

RNA preparation was carried out from 100 mg cell material with the Fast RNA® Pro Blue Kit (MP Biomedicals, Solon, Ohio) following manufacturers instructions. To exclude DNA contamination, the isolated and purified RNA was digested with DNase I for 1 hour at 37°C. Subsequently, the RNA was controlled for DNA contaminations by 18S rDNA PCR. Absence of amplified PCR products was confirmed by agarose gel electrophoresis. The RNA concentration was determined with the Picodrop photometer (Biozym, Oldendorf, Germany).

cDNA synthesis and labeling was done with the superscript II cDNA-Synthesis-Kit according to the manufacturer’s instructions (Invitrogen, Karlsruhe, Germany). A total amount of 10 μg RNA was used for cDNA synthesis.

The labeling of the cDNA was done like recommended by the Nimble Chip Arrays User’s Guide (Gene Expression Analysis v 2.0).

Hybridization of custom 385 K microarray slides with the prepared cDNA was carried out at 42°C overnight using a MAUI Hybridization System following the manufacturer's instructions (BioMicroSystems, Salt Lake City, UT, USA). A stringent washing protocol of the arrays was performed following the instructions of the Nimble Chip Arrays User’s Guide (Gene Expression Analysis v 2.0). The microarrays were stored dry and in the dark until the arrays were scanned.

Chip specifications

The probe sets covering the 5 different bacterial genomes were designed by NimbleGen according to the company’s protocols and algorithms, and all probes were manufactured directly onto the slides via photolithographic synthesis of 60 mer oligonucleotides®. The array probe sets are based on the following whole genome sequences deposited at NCBI: E. nucleatum ATCC 25586 (NCBI, full genome, accession no. AE009931) with 1964 genes, P. gingivalis W83 (NCBI, full genome, accession no. AE015924) with 1883 genes, S. sanguinis SK36 (NCBI, full genome, accession no. CP000387) with 2244 genes, S. mutans UA159 (NCBI, full genome, accession no. AE014133) with 1927 genes, and A. actinomycetemcomitans HK1651 (full genome available at: http://www.oralgen.lanl.gov/oralgen/downloads) with 2168 genes.

For every gene sequence up to 13 individual probes were designed to cover the gene sequence of interest (for detailed information refer to Supporting Tables S3, S5 and S6), resulting in 10,107 unique probe sets covering 10,186 genes. Each probe set was synthesized in three replicates located in different array regions. This setup guaranteed maximum reproducibility of the hybridization process and fluorescence signals. In some cases the NimbleGen algorithms were unable to design adequate probes for certain genes. The genes not covered by the current design are listed in Supporting Table S7.

Additionally, there were 3510 random probes arbitrarily distributed on the chip, which served as negative controls. This entire setup allowed the identification of unspecific reactions without the need to exclude whole genes from the analysis in most cases. The decision if genes with cross-hybridizing probes are eliminated from the investigation could be based on the ratio of nonspecifically reacting probes in relation to the total number of probes for that particular gene. For this study here, we decided to exclude all S. mutans probes which cross-hybridized after S. mitis chip hybridization, but not whole genes from the analysis of the S. mitis/S. mutans two-species combination.

Scanning and quality control of the microarrays and hybridization result interpretation

Scanning of microarrays and subsequent feature extraction was done with the DNA Microarray Scanner from Agilent (Agilent Technologies, Waldbronn, Germany, resolution: 5 μm) and NimbleScan™ software (version 2.4, NimbleGen, Madison, USA), respectively.

Each probe had to pass several criteria of quality control before entering the analysis tool GeneSpring GX (version 11.0, Agilent Technologies, Waldbronn, Germany). These criteria comprised:

i) Identification of probe replicates influenced by wash artifacts: All *pair data files resulting from scanning and feature extraction were analyzed for inhomogeneous replicates of every probe. Replicates with more than a two-fold aberration from the intensity of the other two replicates of this probe were excluded from analysis. In case of more than a two-fold aberration over all three replicates of a single probe, all replicates of this probe were deleted.

ii) Detection of unspecific inter-species cross-reactions: After using cDNAs from all species arranged on the chip for single-species hybridizations, all probes of the other species were checked for their potential hybridization signals. In case of a signal intensity exceeding that of the mean of all random probes on this array by a factor of two, these probes were designated as nonspecifically inter-species cross-reacting (for illustration please refer to supporting Figure S2a–e).

iii) Elimination of results from nonspecifically inter-species cross-reacting probes: When using the cDNAs from the two-species combination S. mutans and S. mitis, cDNA from the latter was checked for cross-hybridization with probes of S. mutans UA159 present on the chip. Employing the criteria from above, a probe was declared as false positive when the derived signal intensity after hybridization with S. mitis cDNA was more than two-fold higher than the mean of all random probes on this array, i.e. the negative control. Again, these nonspecific positive probes were excluded from analysis when comparing the S. mutans single-species transcriptome with the S. mutans transcriptome after co-culture with S. mitis.

Normalization and background correction was done by RMA [Robust Multi-Array Analysis, [40,41]] with the NimbleScan software summarizing all probes per gene resulting in single values per gene. The so obtained *calls data files were analyzed with GeneSpring GX (version 11.0, Agilent Technologies).

All genes of a species, which passed the quality control inspection criteria, were finally grouped into 4 different categories of transcript abundance related to the maximal fluorescence signal intensity of the array (refer to supporting Table S1).

RT-qPCR

Real time PCR (RT-qPCR) was performed for a subset of genes of S. mutans UA159, which were found to be up- or down-regulated in transcriptome analysis of S. mutans UA159 co-cultured with S. mitis compared to the S. mutans mono-species biofilm. Only genes leading to unequivocal hybridization results were used for primer design. Primers were designed with the help of the free online tool Primerfox (http://www.primerfox.com/, Supporting Table S8). For each set of primers, a standard amplification curve was plotted (critical threshold cycle against log of template DNA concentration) and all primers were experimentally controlled for primer
dimer formation. RNA used for cDNA synthesis was controlled for DNA contamination as described earlier in this methods section. cDNA synthesis was done with the superscript II cDNA-synthesis kit from Invitrogen accordant to manufacturer’s instructions.

The RT-qPCR reaction mixture (25 μl) contained 1 × MAX-IMA Sybr Green Master Mix (Fermentas), 6.5 μl cDNA sample and 1 mM of the appropriate forward and reverse PCR primers (Table S3). RT-qPCR reactions were done in 96 well optical reaction plates with the ABI PRISM® 7000 Sequence Detection System (Applied Biosystems). Conditions of RT-qPCR were as follows: 2 min 50 °C, 10 min 95 °C, 15 sec 95 °C, 1 min 60 °C. The last two steps were repeated 40 times.

Each assay was performed with at least three biologically independent RNA samples in technical duplicates.

The critical threshold cycle (Ct) was defined as the cycle in which fluorescence becomes detectable above the background fluorescence and is inversely proportional to the logarithm of the initial number of template molecules. Relative quantification was performed with the comparative Ct method, also known as the 2-ΔΔCt method. Ct values for the genes of interest were normalized to the gene SMU_2015.

Reproducibility and statistics

Each experiment was performed on at least 3 independent occasions (biological replicates) with 2 or 3 replicates each (technical replicates). Statistical parameters (mean, median, standard deviation, p-values) were determined employing the Windows Excel program, and GenSpringGX. P-values less than 0.05 were considered as significant.

Accession number

The raw data and meta information on the experiments have been deposited in the Gene Expression Omnibus database (GEO; www.ncbi.nlm.nih.gov/geo/) and received the accession number GSE28841.

Supporting Information

Figure S1 Documentation of biofilm and monolayer formation of single-species cultures of A. actinomycetemcomitans; F. nucleatum, P. gingivalis, S. sanguinis, and S. mutans. (PDF)

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Figure S2 Histogram of array fluorescence intensities. (PDF)

Table S1 Graduated fluorescence intensities of genes passing the quality control criteria. (XLS)

Table S2 Data set of crosshybridizing probes and the corresponding genes. (XLS)

Table S3 All genes represented on the array, including their appropriate probe number. (XLS)

Table S4 All probes and their corresponding genes which were found as biological false positives after hybridization with RNA of S. mutis. (XLS)

Table S5 All genes represented on the array, with less than 13 probes. (XLS)

Table S6 Genes, which share probes with other genes. (XLS)

Table S7 All genes for which NimbleGen algorithms were not able to identify adequate probes. (XLS)

Table S8 Data set of RT-PCR primer. (PDF)

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Author Contributions

Conceived and designed the experiments: AP BK. Performed the experiments: SR KS. Analyzed the data: SR KS BK. Wrote the paper: SR BK AP KS.

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