ACTINOMYCES NAESLUNDII in initial dental biofilm formation

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The combined use of confocal laser scanning microscopy (CLSM) and fluorescent in situ hybridization (FISH) offers new opportunities for analysis of the spatial relationships and temporal changes of specific members of the microbiota of intact dental biofilms. The purpose of this study was to analyse the patterns of colonization and population dynamics of Actinomyces naeslundii compared to streptococci and other bacteria during the initial 48 h of biofilm formation in the oral cavity. Biofilms developed on standardized glass slabs mounted in intra-oral appliances worn by ten individuals for 6, 12, 24 and 48 h. The biofilms were subsequently labelled with probes against A. naeslundii (ACT476), streptococci (STR405) or all bacteria (EUB338), and were analysed by CLSM. Labelled bacteria were quantified by stereological tools. The results showed a notable increase in the number of streptococci and A. naeslundii over time, with a tendency towards a slower growth rate for A. naeslundii compared with streptococci. A. naeslundii was located mainly in the inner part of the multilayered biofilm, indicating that it is one of the species that attaches directly to the acquired pellicle. The participation of A. naeslundii in the initial stages of dental biofilm formation may have important ecological consequences.

INTRODUCTION

Dental biofilm is an archetypal example of a complex biofilm (Costerton et al., 1999; Davies, 2003; DuPont, 1997). Biofilm formation on tooth surfaces follows the same basic rules as biofilm formation elsewhere in nature. Dental biofilms develop readily because of the optimal temperature, the rich nutrient supply in the oral cavity, and the hard non-shedding surface. They are easily accessible for experimentation using intra-oral devices (Auschill et al., 2002; Palms et al., 2003), and therefore dental biofilms can be used to demonstrate colonization phenomena and ecological principles of universal interest. Concurrent with the increasing recognition of the significance of biofilms in infectious diseases, the development of techniques such as confocal laser scanning microscopy (CLSM), fluorescence in situ hybridization (FISH) and immunofluorescence has enabled visualization of bacteria in their natural undisturbed environment. This offers a substantial improvement upon previous microbiological studies of bacteria grown in planktonic settings (Anwar et al., 1992; Davies, 2003).

Previous studies of dental biofilm that took advantage of these methods mainly focused on streptococci (Diaz et al., 2006; Dige et al., 2007; Hannig et al., 2007; Palmer et al., 2003) because culture-based studies suggested that this group of bacteria is prominent during the initial stages of biofilm formation on teeth (Li et al., 2004; Nyvad & Kilian, 1987, 1990). However, other genera such as Actinomyces are also among the earliest colonizers of dental surfaces and may constitute up to 27% of the pioneer bacteria (Kilian et al., 1979; Li et al., 2004; Nyvad & Kilian, 1987). Several culture-based studies indicated that Actinomyces species gain increased prominence at the expense of streptococci during maturation of the biofilm (Ritz, 1967; Socransky et al., 1977; Syed & Loesche, 1978; van Palenstein Helderman, 1981). Such population changes might reflect differences in growth rates (Nyvad & Kilian, 1987; Socransky et al., 1977) and/or differences in nutritional profiles of these genera (Takahashi et al., 1995; Takahashi & Yamada, 1996; van der Hoeven & van den Kieboom, 1990; Yaling et al., 2006). However, the spatial relationship of actinomyces with other members of the dental biofilm microbiota was not disclosed by these culture-based studies.

Abbreviations: CLSM, confocal laser scanning microscopy; FISH, fluorescent in situ hybridization.

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Previous transmission electron microscopic studies consistently showed the presence of densely packed colonies of pleomorphic Gram-positive bacteria resembling *Actinomyces* in contact with the tooth surface in young and mature supragingival plaque (Listgarten et al., 1975; Nyvad & Kilian, 1987; Schroeder & De Boever, 1970), and similar morphotypes were observed in the demineralized dentine of root surface caries (Nyvad & Fejerskov, 1989). Due to methodological limitations it was not possible at that time to verify the identity of these bacteria. In a recent study using combined CLSM and FISH analysis *Actinomyces naeslundii* constituted up to 18% of the microbiota within the first days of dental biofilm formation, with a notable decrease over the 7 day observation period (Al-Ahmad et al., 2007). Direct imaging of *A. naeslundii* during sequential stages of the initial colonization of hard dental surfaces is still lacking. Such information is important for understanding the role of *A. naeslundii* during initial biofilm development as well as its ecological role in dental disease processes.

The aims of this study were therefore to describe the pattern of colonization and to analyse population dynamics of *A. naeslundii* compared to that of streptococci and other bacteria, during the initial 6–48 h of dental biofilm formation.

**METHODS**

**Experimental conditions.** Oral biofilms were collected on custom-made glass slabs (Menzel). The glass slabs were industrially manufactured (4 × 4 × 1 mm) with a surface roughness of 1200 grit. Six glass slabs were mounted slightly recessed in the buccal flanges of individually designed intra-oral appliances worn by ten healthy volunteers (five females and five males, 23–36 years of age, median age 25 years) for 6, 12, 24 and 48 h. The volunteers retained the appliance intra-orally throughout the experimental period, except during tooth brushing and intake of food or liquids other than water. The Ethics Committee of Aarhus County approved the protocol, and informed consent was obtained from all participants after they received oral and written instructions about the study. A detailed description of the experimental model and the experimental conditions has been previously published (Dige et al., 2007).

**Specimen preparation**

**FISH.** Following *in situ* biofilm growth, FISH was performed as described by Dige et al. (2007), using specific 16S rRNA probes against streptococci, *A. naeslundii* and all bacteria. Immediately after removal from the oral cavity, the glass slabs with the biofilms were fixed in 4% paraformaldehyde (3 vols) in PBS (1 vol.) (Stahl & Amaril, 1991) for 3 h at 4 °C. The specimens were subsequently washed with sterile PBS and stored in a mixture of 100% ethanol and PBS (1:1) at −20 °C. For permeabilization, the glass slabs with the biofilm were mounted on diagnostic glass microscope slides (Menzel) with paraffin wax (GC Corporation) and treated with 25 μl lysozyme (Sigma) [70 U μl⁻¹] in 100 mM Tris/HCl pH 7.5 (Sigma), 5 mM EDTA (Merck)] for 9 min at 37 °C in a humid chamber. The diagnostic glass microscope slides with the biofilms were then rinsed with ultrafiltered water, dehydrated in series of ethanol washes (50, 80 and 100%; 3 min each wash) and dried for 10 min in a vertical position. The glass slabs were then exposed to 10 μl hybridization buffer (0.9 M NaCl, 20 mM Tris/HCl pH 7.5, 0.01% SDS, 30% formamidine, which pilot studies showed to be the optimal concentration for the probe/label combination used) containing 100 ng of the designated oligonucleotide probe and incubated at 46 °C for 2 h in a humid atmosphere in the dark. After hybridization, the glass microscope slides were first washed in buffer (20 mM Tris/HCl pH 7.5, 5 mM EDTA, 0.01% SDS and 112 mM NaCl) for 15 min in a water bath at 48 °C, and then rinsed in ice-cold ultrafiltered water. The oligonucleotide probe STR405 (5'-TAG CCG TCC CTT TCT GGT-3') (MWG Biotech) labelled with Alexa488 was used to identify all Streptococcus spp. (Paster et al., 1998) and the oligonucleotide probe ACT476 (5'-ATC CAG CTA CCG TCA ACC-3') (IBA) labelled with Atto550 was used to identify *A. naeslundii* (Gmür & Lüthi-Schaller, 2007). The oligonucleotide probe EUB338 (5'-GCT GCC TCC CGT AGT AGT-3') (IBA) labelled with Atto635 was used as a positive control based on its ability to detect all bacteria (Amann et al., 1990) with a few exceptions such as Treponema multilopithium and Treponema lecitinolyticum (Daims et al., 1999), which are not involved in the early phase of biofilm formation. A search performed in the Ribosomal Database Project II at http://rdp.cme.msu.edu/index.jsp indicated that the probe EUB338 recognizes 241,803 out of 335,830 bacterial sequences in the database, including all taxa of bacteria hitherto detected in the oral cavity.

sequence in the positive species and single to multiple sequence deviations in the negative species (Fig. 1). Based on these observations it was concluded that probe ACT476 reacts only with *A. naeslundii* among species that are associated with humans. The DAPI reagent stained all bacteria in the panel.

**CLSM.** The glass slabs with the biofilm were examined on an inverted Zeiss LSM 510 META confocal microscope using 488, 543 and 633 nm laser lines for excitation of Alexa488, Atto550 and Atto633, respectively. Emission bandpass filters were set to 500–530 nm for detection of Alexa488, 565–615 nm for detection of Atto550, and 651–704 nm for detection of Atto633. Images were acquired using a ×63 (NA=1.2) water-immersible objective (C-Apochromat) with a working distance of 0.28 mm and collected at electronic zoom ×0.7 (2048×2048 pixels, pixel size 100 nm=optimum resolution), ×2 (1024×1024 pixels, pixel size 70 nm) and ×4 (1024×1024 pixels, pixel size 30 nm), of which the last was used for quantification. Images were acquired with pinholes set to 1 Airy unit, corresponding to an optical slice thickness of 1.0 μm. Prior to microscopy the glass slabs with the biofilms were mounted on a plastic device so that the biofilm could be turned upside down in a chambered coverglass for examination by the inverted microscope. A drop of Citifluor AF3 anti-bleaching reagent was applied between the biofilm and the chambered coverglass. For the qualitative analysis each biofilm was scanned at representative areas, avoiding the edges of the glass slabs. Two series of optical sections were generated by vertical sectioning at 1 μm distances through the biofilm. Because the surface was never exactly horizontal with respect to the microscope stage, the number of optical sections per specimen was not directly translatable to biofilm thickness. For quantitative analysis (zoom 4) each biofilm was scanned using a systematic uniformly random sampling method (see description of the stereological analysis), and z series of optical sections were generated by vertical sectioning at 0.5 μm distances through the biofilm. This interval was chosen in order to include the major bacterial groups of supragingival biofilm.

**Quantification of bacteria.** The number *Q* of bacteria was counted using the unbiased counting frame originally described by Gundersen (1977) and applied to bacteria, as described by Dige et al. (2009). The unbiased counting frame was superimposed on the images and fixed in the same position throughout subsequent focal planes. Bacteria present was identified as a 2 m·m² quadrant in the centre of each glass slab in order to capture typical smooth surface biofilm. Subsequently, based on a visual inspection of the density of bacteria, four or eight systematic uniformly random sampling fields were chosen. The first field of view was sampled using a random number table. From this random starting point within the area of interest, the remaining three or seven fields of view were sampled by moving the microscope stage with a fixed distance from the previous field (in this case 1000 μm in the x-axis and 500 or 1000 in the y-axis, depending on whether four or eight sampling fields were chosen). The principle of counting has been previously described (Dige et al., 2009).

**Image analysis.** Image analysis was performed using ImageJ 1.34s (Abramoff & Viergever, 2002; Rasband, 1997–2006). ImageJ was used to adjust output levels within the individual channels of the 24-bit RGB merged images. Prior to merging, the images for each colour channel were assembled into image stacks. In the merged images, streptococci, *A. naeslundii* and remaining bacteria were represented by green (yellow), blue (purple) and red colours, respectively. No other manipulation of the images was performed. For illustration purposes, maximum projection images of the entire confocal image stack were made for some 6 h, 12 h and 24 h specimens to compensate for the glass surfaces not being oriented completely parallel to the optical section plane.

**Stereological analysis**

**Systematic uniformly random sampling.** Stereological analysis was performed as a systematic uniformly random sampling of fields of view (Gundersen & Jensen, 1987) as previously described by Dige et al. (2009). First, the area of interest for estimating the number of bacteria present was identified as a 2 × 2 mm² quadrant in the centre of each glass slab in order to capture typical smooth surface biofilm. Subsequently, based on a visual inspection of the density of bacteria, four or eight systematic uniformly random sampling fields were chosen. The first field of view was sampled using a random number table. From this random starting point within the area of interest, the remaining three or seven fields of view were sampled by moving the microscope stage with a fixed x and y distance from the previous field (in this case 1000 μm in the x-axis and 500 or 1000 in the y-axis, depending on whether four or eight sampling fields were chosen). The principle of counting has been previously described (Dige et al., 2009).

**Fig. 1.** Comparison of the target sequence of probe ACT476 in *Actinomyces* species shows that reactivity of the probe is restricted to members of the perfectly matching *A. naeslundii–A. viscosus–A. bowdeni–A. denticolens* cluster, among which only *A. naeslundii* is associated with humans. The phylogenetic tree of *Actinomyces* species shown to the left was based on complete 16S rRNA gene sequences clustered by the Minimum Evolution algorithm in MEGA version 4 (Tamura et al., 2007).
were only counted the first time they came into focus in a section. Bacteria were counted manually, and to remember which bacteria had already been counted, the point picker in the Particle Analysis plugin in the ImageJ software was used. The software also maintained a record of the number of cell markers placed by the operator. Because of the pleomorphic morphology of ACT476-labelled bacteria the following counting rules were adopted: (i) for overlapping bacteria (intense fluorescence) the bacteria were counted as separate bacteria; (ii) when a space or a notch was observed, bacteria were counted as separate bacteria; (iii) bacteria showing a change in angle were counted as separate bacteria (Fig. 2). ACT476-labelled bacteria were counted on images of the blue channel only (with the other channels off), because of better differentiation when they were not intermingled with other types of bacteria (compare Fig. 4g and h). The principle for counting streptococci in division was to count them as two bacteria when the length was equal to that of two separate cocci.

The size and number of the unbiased counting frame varied according to cell density. For example, in specimens with low cell density (6 and 12 h), one counting frame was used covering the whole image except the outermost borders. On the other hand, when the cell density was high (multilayered biofilms) four counting frames of smaller areas were superimposed on the images. The strategy was such that for each specimen, approximately 100–200 bacteria needed to be counted to get reliable quantitative data (Gundersen et al., 1999; Nyengaard, 1999). The dimensions of the counting frame(s) remained the same for all four/eight images in one specimen.

The raw counts of streptococci, A. naeslundii and remaining bacteria were used to estimate the total bacterial number within the area of interest (2 × 2 mm²). Subsequently, the total number of bacteria for each glass slab, N, was estimated by the 2D fractionator (Dige et al., 2009; Gundersen, 1986). For each volunteer two glass slabs were analysed at each time point and the mean value of the estimates was calculated.

Statistics. For each bacterial group the number of bacteria was plotted as a function of time on log-linear and log-log scales and evaluated for linear behaviour, signifying single-exponential or nonlinear growth, respectively.

The total variation between individuals (CVtot) was determined, as regards the number of streptococci, the number of A. naeslundii and the total number of bacteria. The error variance due to the stereological method (CEmet) was estimated as the counting noise (Nyengaard, 1999), disregarding the error variance due to systematic sampling of sections and fields of view. The observed total variation [CVtot=standard deviation (SD) divided by the mean] was calculated. From the CVtot and the CEmet, the biological variation, CVbio, was determined using the equation CVbio=CVtot+CEmet. Because the error variance due to the stereological method (CEmet) was very small (Dige et al., 2009) the total variation between individuals (CVtot) in the number of bacteria, as regards streptococci, A. naeslundii and total bacteria, was determined mainly by the biological variation (CVbio). Calculation of the ratios of CE2/CV2 gave very small values, suggesting that a sufficient number of bacteria was counted (Nyengaard, 1999).

RESULTS

Quantitative observations

Fig. 3(a) shows the number of streptococci (green) and the number of A. naeslundii (blue) in biofilms developed during 6, 12, 24 and 48 h from the ten individuals. The distribution of the number of streptococci and A. naeslundii for the ten individuals is further illustrated in Supplementary Fig. S1 (available with the online version of this paper) by pairs of data with coloured lines. The speed of bacterial surface coverage varied considerably between individuals at all time points, but for both bacterial groups there was a notable increase in the number of bacteria over the observation period. A plot (Fig. 3b) of the slopes for the increase in ln(number of streptococci) versus the slopes for the increase in ln(number of A. naeslundii), evaluated as a function of increase in ln(time), showed a tendency towards a faster increase for streptococci than for A. naeslundii, although this was not statistically significant (paired t-test, P=0.07). This trend was supported by an analysis of changes in the proportion of bacteria between 6 and 48 h, which showed that in nine of the ten individuals there was a tendency to an increase in the proportion of streptococci relative to total bacteria, whereas the proportion of A. naeslundii decreased slightly or remained stable in nine of the ten individuals (Fig. 3c, d).

Qualitative observations

At 6 and 12 h, bacteria were scattered randomly across the surface, as single bacteria or as mono- and multi-genera

Fig. 2. Illustration of the principle of counting ACT476-labelled bacteria. The figure shows four consecutive x-y sections of 6 h biofilm on images of the blue channel only (with the other channels off), demonstrating A. naeslundii (blue). Bacteria were only counted the first time they came into focus in a section, as indicated by white * in the consecutive sections. For detailed explanation of the counting principle, see text. Scale bar, 5 μm.
clusters of bacteria, many of which appeared in a stage of cell division (Fig. 4a–e). In all individuals *A. naeslundii* was observed at all time points scattered throughout the biofilm. In the early stages *A. naeslundii* was in all individuals recorded as single coccoid rods or short rods arranged in Y, V and T shapes (Fig. 4a, b, d), and in six of the ten individuals also as longer rods or filamentous bacteria (Fig. 4a, c, e). *A. naeslundii* was observed both as isolated clusters (Fig. 4a, c) and in mixed clusters with streptococci and/or other bacteria (Fig. 4a, d, e). Notable differences in the amount of bacteria and the composition of the microbiota were observed both across the experimental surface and between surfaces carried by different individuals.

At 24 h and 48 h, the biofilm showed dominance of streptococci (Figs 4f, g and 5b). Three individuals, in addition, showed large accumulations of non-streptococci (Fig. 5a), including *A. naeslundii* and large coccoid non-streptococci in pairs or tetrads. Eight of the ten individuals showed *A. naeslundii* arranged in microcolonies of varying size consisting of branching filaments, some of which were ‘spider colonies’ consisting of branching filaments radiating from a single point (Fig. 4f, g, h). *A. naeslundii* was also observed intermingling with streptococci and other non-streptococci in most individuals (Fig. 4f, g). Also at these more advanced stages of biofilm formation the pattern and degree of microbial coverage, as well as the thickness of the biofilm, varied within and between individuals from incomplete (Fig. 4f) to complete surface coverage by bacteria (Fig. 4g), in some parts with prominences (‘chimneys’) of multilayered complex microcolonies (Figs 5 and 6). All types of bacteria showed various stages of cell division reflected by their pair-wise (Fig. 4f) or branching arrangement (Fig. 4f, h).

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**Fig. 3.** (a) The number of *Streptococcus* spp. (green) (Dige et al., 2009) and the number of *A. naeslundii* (blue) recorded in dental biofilms developed within 6, 12, 24 and 48 h in ten individuals. Note the logarithmic scale on the y-axis. The estimated numbers refer to a well-defined reference space that corresponds to an area of 4 mm². (b) The relationship between the slopes for the increase in number of streptococci versus the slopes for the increase in ln(number of *A. naeslundii*), evaluated as a function of the increase in ln(time), showing a tendency towards a faster increase for streptococci than for *A. naeslundii*. However, this was not statistically significant (paired t-test, *P*=0.07). The bold line indicates an equal slope for the increase in number of *A. naeslundii* and streptococci. (c, d) Linear association (trend lines) of the changes in the proportion of *Streptococcus* spp. (c) and *A. naeslundii* (d) as a percentage of the total number of bacteria between 6 and 48 h in ten individuals. Note different scales on the y-axis in (c) and (d).
Examination of the biofilms in the z-axis (x-y sections) allowed a detailed analysis of the central parts of the ‘chimneys’. The central parts of the ‘chimneys’ varied in composition and were either sparsely colonized (Figs 5b and 6b) or composed of non-streptococci (Figs 5a and 6a), including large coccoid bacteria, *A. naeslundii*, other rod-shaped/filamentous bacteria, or other non-streptococci and non-*A. naeslundii*. Analyses along the z-axis further demonstrated that, at the surface of the multilayered biofilms, bacteria of different genera regularly co-localized.

Analysis of consecutive sections of the multilayered biofilm parallel to the surface showed that *A. naeslundii* predominantly colonized the inner part next to the glass surface, and was more sparsely distributed in the outer layers (Fig. 5). Some *A. naeslundii* microcolonies extended perpendicularly from the supporting surface surrounded by other bacteria and forming ‘chimney structures’. Sagittal (x-z, y-z) sections confirmed the presence of *A. naeslundii* in the inner layers as individual pleomorphic bacteria were oriented at different angles to the surface and forming palisades (Fig. 6).
DISCUSSION

Several Actinomyces species belong to the resident oral microbiota of supra-gingival plaque, although studies based on culture, checkerboard hybridization, 16S rRNA gene libraries and FISH reveal significant differences in their proportions depending on the age of the biofilm (Al-Ahmad et al., 2007; Diaz et al., 2006; Haffajee et al., 2008; Li et al., 2004; Ramberg et al., 2003). Using a species-specific oligonucleotide probe this study confirms the checkerboard hybridization-based demonstration of A. naeslundii as a significant member of the initial colonizers of tooth surfaces. For the first time, our study provides a presentation of the spatio-temporal organization of A. naeslundii in relation to other bacteria in initial multi-layer dental biofilms formed in vivo up to 48 h and definitively demonstrates that A. naeslundii preferentially occupies the inner part of early multilayered biofilms. We therefore infer that the densely packed, pleomorphic Gram-positive bacteria with thick cell walls previously observed close to the enamel surface in electron micrographs (Listgarten et al., 1975; Nyvad & Fejerskov, 1987b; Schroeder & De Boever, 1970) are A. naeslundii.

A. naeslundii was often observed in mixed clusters with streptococci and other bacteria at 6 and 12 h. This observation supports the view that co-adhesion, in particular co-adhesion processes involving A. naeslundii, streptococci and other bacteria, play an important role during the initial stages of colonization of tooth surfaces (Bos et al., 1996; Gibbons & Nygaard, 1970; Kolenbrander, 1988; Kolenbrander et al., 1990; Palmer et al., 2003; Yoshida et al., 2006). This observation is further supported by the finding of genotypically different bacteria co-localizing at the outer surface of the biofilm, indicating that co-adhesion of bacteria from saliva is a continuing process adding to the biomass of the developing biofilm. However, it is conceivable that cell division is the major contributor to the rapid increase in biomass during the first 24–48 h of biofilm formation, as suggested by several reports (Bloomquist et al., 1996; Skopek et al., 1993; Ørstavik, 1984). This is in line with the results of the present study, in which many bacteria appeared in a stage of cell division, including A. naeslundii, which formed branching filaments or ‘spider colonies’. Likewise, it is conceivable that the decrease in the relative proportion of A. naeslundii between 6 and 48 h of biofilm formation observed in this and previous studies (Al-Ahmad et al., 2007; Li et al., 2004) reflects the effect of cell division and is a direct result of slower cell division of A. naeslundii relative to streptococci and other members of the microbiota. Similar shifts in the relative composition of the microbiota have been recorded during biofilm formation in experimental rats (Beckers & van der Hoeven, 1984).

The observation of densely packed colonies of A. naeslundii in the innermost part of the biofilm adjacent to the supporting surface has interesting ecological implications.

Fig. 5. CLSM images of 48 h in situ biofilm showing consecutive x-y sections at 4 μm intervals (along the z-axis) from the supporting glass surface (bottom image) in two different individuals. Biofilms were stained simultaneously with all-bacterium-specific EUB338 probe, Streptococcus-specific STR405 probe and Actinomyces-specific ACT476 probe. Green (yellow), blue (purple) and red represent streptococci, A. naeslundii and the remaining bacteria, respectively. A. naeslundii (purple-blue) are predominantly located in the inner part next to the glass surface, whereas these species are more sparsely distributed in the outer layers. In both illustrations columnar microcolonies protrude as circular projections (chimneys) at varying distances from the supporting surface. In (a) the central part of the microcolonies is composed of A. naeslundii (arrows) or other non-streptococci whereas in (b) the central parts are often sparsely colonized (arrows). Scale bar for all images, 50 μm.
In contrast to streptococci, *A. naeslundii* has a unique glycolytic system in which the bacteria use phosphoryl donors instead of ATP for carbohydrate degradation (Takahashi *et al.*, 1995). *Actinomyces* species can use lactate as a carbon source for growth (Takahashi & Yamada, 1996; van der Hoeven & van den Kieboom, 1990), whereby lactic acid is converted into weaker acids (Takahashi & Yamada, 1996). A pH-modulating activity of these species may, theoretically, occur also via degradation of urea (Yaling *et al.*, 2006). Moreover, through its metabolism, *Actinomyces* species can remove oxygen from the environment and create an anaerobic milieu (Takahashi & Yamada, 1996), suitable for outgrowth of some other bacteria. Finally, recent observations demonstrate that co-aggregation with *A. naeslundii* stabilizes arginine metabolism in *Streptococcus gordonii* and reduces its dependence on extracellular arginine, which is a limiting factor in the environment of the early colonizers (Jakubovics *et al.*, 2008; Van Wuyckhuyse *et al.*, 1995). Collectively, these properties make *A. naeslundii* an essential initial colonizer of tooth surfaces and particularly well adapted to live and survive in substrate-limited environments deep in the biofilm. The concerted metabolic activities of these bacteria may have a controlling effect on dental caries processes by reducing the acidogenic potential of the biofilm (Takahashi & Nyvad, 2008).

With increasing age of the biofilm, microcolonies of *A. naeslundii* and other non-streptococci were seen to extend perpendicularly from the supporting surface as ‘chimney’ structures and palisades like those observed by electron microscopy of multi-layered dental plaque (Listgarten *et al.*, 1975; Nyvad & Fejerskov, 1987b; Rosan *et al.*, 1976). The morphogenesis of these particular structures can only be speculated on. It may reflect a constrained physical environment during development of the biofilm whereby overgrowth of rapidly multiplying species may hinder the growth of other bacteria with a lower growth rate such as *Actinomyces* species. Alternatively or additionally, such structures may result from nutritional interrelationships between different microbial species or specific co-adhesion/co-aggregation processes. Thus, Bos *et al.* (1996) proposed that streptococci may encapsulate *Actinomyces* to form micro-anaerobic domains in the biofilm, which are needed for optimal growth of the *Actinomyces*. This hypothesis corroborates more recent concepts of bacterial multicellularity that bacteria growing in biofilm communities have communication and decision-making capabilities that enable them to coordinate growth and biochemical activities (Jakubovics *et al.*, 2008; for reviews see Kolenbrander *et al.*, 2006; Shapiro, 1998). Hence, it has been suggested that the growth rate of adherent cells is enhanced when a certain cell density is reached, whereas the growth rate drops at higher densities. This density-dependent growth may be explained by cell–cell signalling, resulting in physical or morphological changes of the biofilm bacteria (Bloomquist *et al.*, 1996).

In this study, *A. naeslundii* represented a large spectrum of morphotypes, ranging from coccoid to small rods and filamentous bacteria. It has been suggested previously that *A. naeslundii* exhibits pleomorphism, the coccoid form predominating during the early stages, whereas rod-shaped or filamentous forms become prominent after 24–48 h (Nyvad & Fejerskov, 1987b). This observation is consistent with our study (compare Fig. 4b with Fig. 4f, h) as well as immunoelectron microscopic studies of dental plaque *in situ*, in which *A. viscosus* (*A. naeslundii* according to present nomenclature) tended to be coccobacillary in the superficial layers and filamentous in the deeper layers (Berthold *et al.*, 1982).

The observation of sparsely colonized areas in the centre of the circular projections and deeper parts of multilayered biofilms (Fig. 5) is open for speculation. Such unstained regions have been suggested to represent open voids (Pratten *et al.*, 2000; Wood *et al.*, 2000) or to contain extracellular polysaccharides (Thurnheer *et al.*, 2004). However, one cannot exclude the possibility that these areas contain bacteria labelled with dyes that are bleached away by out-of-focus excitation during the consecutive scanning through the biofilm, or bacteria exhibiting

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**Fig. 6.** CLSM images of 48 h *in situ* biofilm showing sagittal (x-z, y-z) sections of specimens from each of the two individuals shown in Fig. 5. Note that *A. naeslundii* (blue) are predominantly identified in the inner part next to the glass surface. Some microcolonies of *A. naeslundii* extended almost throughout the entire thickness of the biofilm (a). Note also sparsely colonized zones next to the supporting surface (b). The width of both images is 200 µm, and the heights of the images are 18 µm and 23 µm, respectively.
insufficient fluorescent signals because of low rRNA content due to slow growth or low metabolic state (Ammann et al., 1995; Hannig et al., 2007; Moter & Göbel, 2000; Schupper et al., 1998). In fact, in some instances we found the fluorescent signal of the ACT476 probe to be bright, whereas the signal of the EUB338 probe of the same bacteria was very low or absent. Of particular relevance to our stereological approach, it has been previously observed that the in situ hybridization of Gram-positive filamentous bacteria such as Actinomyces often results in an irregular distribution of fluorescent signals over the whole filaments (Schupper et al., 1998), possibly because of insufficient permeability of the bacterial cell walls, which has been documented also for actinomyces in other ecosystems (Müller et al., 2007; Schupper et al., 1998). Consequently underestimation of the number of some bacteria cannot be excluded (Dige et al., 2009).

In conclusion, by combining qualitative and quantitative methods this study resulted in new insight into the temporo-spatial relationships as well as the population dynamics of A. naeslundii relative to streptococci in the initial phases of biofilm formation on oral solid non-shedding surfaces. A remarkable observation of the study was the preferential colonization of A. naeslundii in the deeper regions of the biofilm. In view of the pH-modulating properties of A. naeslundii it is relevant to further explore the ecological role of this species in the processes of dental caries.

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