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Title	Factors involved in Candida biofilm formation on acrylic surfaces
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Periodontopathogenic Bacterial Biofilms and Planktonic Cells - An Ultrastructural Comparison N MORDAN* D JENSEN, M WILSON, and H NEWMAN. Eastman Dental Institute, London, UK. Dental plaque, the actiologic agent of both dental caries and chronic inflammatory periodontal disease, is a bacterial biofilm formed by adherence of planktonic free-living bacteria to a surface and their subsequent proliferation. Much past research has involved the assessment of bacterial death in planktonic culture with a view to arresting the advance of plaque <i>in vivo</i> . However it is now apparent that the behaviour of members of a biofilm, in particular their resilience to adverse conditions differs from that of planktonic cells. In this study we cultured <i>Actinobacillus actinomycetemcomitans</i> in Wilkins Chalgren both with 10% defibrinated horse blood and as a biofilm (on cellulose nitrate membrane filters on Wilkins Chalgren agar plates) for 24, 48 and 72 hours. Resulting cultures were fixationed with glutaraldehyde, post-fixed in osmium tetroxide, dehydrated and embedded in LR White resin Ultrathin sections were stained with uranyl acetate and lead citrate and viewed in a JEOL 100CX transmission electron microscope. Although there was progressive alteration in the appearance of both protoplasm and cell wall with time, the rate of change was faster for cells in broth as opposed to biofilm. Planktonic bacteria showed intact cells at 24hrs but by 72hrs there was advanced lysis and protoplasmic condensation. However biofilm cells displayed only slight lysis and little protoplasmic condensation at 72hrs. These differences in ultrastructure between planktonic and biofilm bacteria are evidence of the slower growth rate of cells in biofilm which in turn may account for the refractory nature of biofilms.	128500 Restance of <i>Porphyromonas gingradis</i> in biofilms to doxycycline and metonidazole T LARSEN* and B. CIWERCMAN (Institute of Odontology, Eaculy of Health Sciences, University of Copenhagen, Copenhagen, Denmark). The ways to determine the susceptibility of <i>Porphyromonas gingradis</i> (<i>Pg</i>) in biofilms of the present of doxycycline and metronidazole. 48h biofilms of three reference strains and three clinical isolates of portion of the strains in the strain of the present of doxycycline and metronidazole. 48h biofilms of the reference strains and three clinical isolates of portion of the strains in the strain of the present of planktonic bacteria by rinsing, the filters were transferred to tryptic soy agar plates (TSA, Difco) containing doxycycline of dyng-free TSA for determination of the minimal bactericidal concentration (MEC). It is and the clinical isolates of planktonic bacteria of the same strain and a reference strain were inoculated, too. After incubation for 7d non-growing bacteria were transferred to dyng-free TSA for determination of the minimal bactericidal concentration (MBC). The MBC of metronidazole the MBC of doxycycline to biofilm bacteria increased up to more than 100 of planktonic bacteria in the MBC of doxycycline to biofilm bacteria increased to biofilm the the MBC of doxycycline to biofilm tacteria increased to biofilm tells while a minor increase was enfor planktonic cells. The MBC of metronidazole to biofilm MIC-value. The MBC of doxycycline and metronidazole to biofilm MIC value. The MBC of doxycycline and metronidazole to biofilm tells while a minor increase was sent for planktonic bacteria to reased to a lamost increased to a the MBC of doxycycline and metronidazole to biofilm tells while a minor increase was sent for planktonic cells. The MBC of metronidazole to biofilm tells while a minor increase was sent for planktonic bacteria to reased to a lamost intereased, too the MBC of doxycycline to biofilm cells with the materia increased to the metronidazole to biofilm tells increase
2857 Community structure and enzyme activity in microcosm dental plaques. R.J. PALMER JR.*. L. WONG' and C.H. SISSONS' (Ctr. Env. Biotech., University of Tennessee, Knoxville; 'Dental Research Group, Wellington Sch. Med., Otago University, Brance activity in <i>vitro</i> microcosm dental plaques. Microcosm plaque biolifuns were cultured in a multi-plaque antificial mouth from a plaque-enriched salivary inoculum under continuous supply of a saliva-analogue solution that contained protein in four different hydrolysis states: 0.25% mucin analogue solution that contained protein in four different hydrolysis states: 0.25% mucin and a mixture of amino acids equivalent to 5 g/L of casein plus 0.25% mucin, Sg/L of a partial digest of casein (rhypticase Peptone) plus 0.25% mucin, and a mixture of amino acids equivalent to 5 g/L of casein plus 0.25% mucin. These former communities also had fewer lipid-based generic groups (4) than did the latter (5 in the approxemution control supplied plaque; 6 in the amino-acid-supplied plaque). Capnocytophaga occurred only in the amino-acid supplied plaque, and Bacteroides was found in higher amounts in plaques supplied with peptone or with amino acids than in those supplied with unhydrolyzed protein; a more Gram negative and complex microflora occurred when simplied when unhydrolyzed protein; a sing API-ZYM™ plates showed that in plaques grown on mucin alone, slightly higher levels of cysteine arylamidase and C16 lipase were present, but lower levels (2: to 4-fold) of 15 other enzyme activities occurred: complex unbixrates seemed to decrease erzymatic (metabolic) diversity. We conclude that community structure and whole singliate are stronely dependent on environmental lactors such as the degree of hydrolyzed protein; a protein on environmental lactors such as the degree of hydrolyzed prot	2852 Topographical vitality pattern in dental plaque biofilms AUSCHILL, T.*, NETUSCHIL, L., BRECX, M., SCULEAN, A., REICH, E. (Periodontology & Conservative Dentistry, Homburg/Saarland, Germany): A vital fluorescence technique was combined with optical planes analysis using a confocal laser scanning microscope (CLSM). By combining these techniques the three-dimensional vitality distribution in plaque biofilms of a light (LPF) and a heavy plaque former (HPF) was evaluated. Both volunteers wore enamel chips in a splint for five days. After vital staining with fluoresceine diacetate and ethidium bromide the specimens were processed for the CLSM examination. One μ m thick optical planes were analyzed in the z-axis of these dental plaque biofilms. The plaque biofilm of the LPF was 16 μ m high, sparse and showed low vitality, <i>i.e.</i> < 10%. On the other hand, that of the HPF was heigher (26 μ m) and more vital, <i>i.e.</i> up to 30 %. In both instances the bacterial vitality increased from the enamel surface to the external part of the dental plaque. However, these overall data do not explain the spatial distribution of the vitality. Looking closely, it was found that voids in the plaque biofilm structure were outlined by layers of vital bacteria, while these were packed into layers of dead material. More CLSM-data of undisturbed plaque biofilms are therefore needed to reveal the complex three-dimensional adhesion and growth pattern of human dental plaque. This study was supported by GABA International, Basel, Switzerland.
2853 Artificial Bacterial Biofilms and the Confocal Laser Scanning Microscopic Analysis. S.TAKENAKA*, M. IWAKU and E. HOSHINO (Operative Dentistry and Endodontics and Oral Microbiology, Niigata University School of Dentistry, Niigata Japan). Bacterial biofilms may be formed at various sites, including on mucos membrane, teeth and even in infectious lesions. To elucidate the structure and the function of biofilms, mucoid-type Pseudomonas aeruginosa organisms (strain PT1252) were centrifuged onto surface of a cover-glass and cultured in broth media supplied continuously (45 ml/hr). The biofilm structure of 4, 6, 8, 12 and 24 hr-old was visualized by fluorescent staining (SYTO9, PI and/or FITC-ConA). It was clearly demonstrated that No. of bacteria (10°-10°) can be estimated by the fluorescent intensity. Counting of each biofilm layer (1µm depth) revealed clearly that the biofilm developed 3-dimensionally to make up 9 layers after 12 hrs. The living and dead organisms were differentiated by SYTO9 and PI, respectively, <i>in situ</i> in biofilm, and about 10% organisms were kalled, but 11±1.3% (n=3) of organisms in 24 hr-old biofilm were still alive after the exposure for 120 min. These results indicate that the confocal laser scanning microscopical analysis of artificial biofilms was useful to elucidate bacterial functions in biofilms, and may lead to new quantitative estimation of bactericidal efficacy of antibacterial drugs.	2854 Detection of pH Gradients in Biofilms Using 2-Photon Excitation Microscopy, D.J. BRADSHAWY ¹ , P.D. MARSH ¹ , H. GERRITSEN ² , J. VROOM ² , G.K. WATSON ³ & C. ALLISON ³ (CAMR ² , Salisbury, UK, University of Utrecht ² , Utrecht, The Netherlands & Univer Research ³ , Bebington, UK). Gradients in key environmental parameters develop in biofilms. Defined, mixed culture biofilms of oral bacteria have been generated in a chemostat model (<i>J Appl Bacteriol</i> 80, 124-130, 1996). These biofilms were reproducible, but only 5-20 µm in depth, whereas plaque associated with fissures, or at the gingival crevice, is far deeper. This study aimed to produce biofilms of between 100-400 µm depth, and to examine these biofilms using a 2-photon excitation fluorescence lifetime microscope; pH gracients within the biofilms were measured directly using a pH-sensitive dye. A constant depth film fermenter (CDFF) was inoculated for 8 hours at 50 ml.h ⁻¹ with mixed cultures of up to 10 species of oral bacteria grown in a chemostat, fresh growth medium was supplied to the biofilms was determined using selective and non-selective agar media. Replicate biofilms were everial with 2% (w/v) sucrose for 1 hour, and then carboxyfluorescen inded. The distribution of pH was then determined by examining fluorescence lifetime images throughout the depth of the biofilm. Viable counts showed that diverse mixed culture biofilms developed, including aerobic, faculative and anaerobic species. The 2-photon excitation microscope was able to distinguish individual bacteria to a depth of up to 140 µm. The sucross/carboxyfluoresceni images indicated zones with pH as low as pH 30, in some cases adjacent to areas with much higher pH (pH > 5.0). The CDFF apparatus can generate deep biofilms to simulate plaque from at-risk sites. The 2-photon excitation microscope has been used successfully to resolve images as deep as 140 µm, and allowed the real-time imaging of pH in biofilms.
Effect of endogenous proteins on Candida biofilm formation on acrylic H. Yamashiro, H. Nikawa, T. Hamada, H. Nishimura and L.P. Samaranyake ¹ (Hiroshima University, Hiroshima, Japan, ¹ Hong Kong University, Hong Kong, China) It is known that fibronectin(FN), mannan-binding proteins(MBP), mucin and concanavalin-A(Con-A) binding material may be involved in candidal biofilm formation on acrylic surfaces (Nikawa et al. Microbial Ecol Health Dis 9: 32-43, 1996). These proteinances as are derived in vivo from endogenous sources such saliva, serum and other microbial products. In the current study we further examined, using the method of Nikawa et al (1996) the effect of salivary or serum proteins, such as mucin, fibronectin and MBP, on in vitro candidal biofilm formation. A single oral isolate each of <i>Candida abicans, C. torojacidis and C. glabaratic were used in the study.</i> The biofilm activity on acrylic strips laced with the appropriate proteins and the yeasts was determined using the previously described adenosine triphosphate (ATP) assay. The biofilm activity was assessed after supplementation of human saliva/serum with various proteins. Supplementation of human mixed saliva with FN had no significant effect on biofilm activity whereas supplementation of human mixed saliva with FN had no significant exting of <i>C. glabarat</i> as compared with the controls. In contrast, the solution may not support the biofilm formation, of <i>Candida</i> isolate salivary mucin and FN la isolation may not support the biofilm formation, of <i>Candida</i> isolate (pc0.01). This observation implies that FN together with protect biofilm activity of all Candida isolates (pc0.01). This observation implies that FN together with biofilm activity of <i>C. albicanstropicalis</i> (pc0.03). Thus it seems that the cell surfacently reduce the biofilm activity of <i>C. albicanstropicalis</i> (pc0.03). Thus it seems that the cell surfacently of the the studies to the surgest that a mubber of formadigenous proteins present both in saliva and serum contribute to <i>Can</i>	2856 Factors involved in <i>Candida</i> biofilm formation on acrylic surfaces H. Nikawa, T. Hamada, H. Yamashiro and L.P. Samaranayake ¹ (Hiroshima University, Iiroshima, Japan, ¹ Hong Kong University, Hong Kong, China) Despite the realization that successful colonization of denture surface is an important step in the denture stomatitis, the role played by denture pellicle during the colonization process and subsequent biofilm formation is poorly understood. In addition, it is likely that during biofilm formation, cell surface properties of <i>Candida</i> , may be involved and thus a deeper understanding <i>Candida</i> biofilm formation necessitates clarification of such intertwined relationships. The purpose of the present study was to analyze fungal properties, such as acid production, cell surface hydrophobicity, germ tube formation and thigmotropism, involved in the <i>Candida</i> biofilm formation on saliva, serum, and saliva- serum coated asrylic determined by ATP (adenosine triphosphate) analysis. The biofilm activity ouried depending upon both the isolate and the pellicle, As compared with the suscoated cortrol, significantly increased biofilm activity of the protein-free control strips, with all isolates of <i>C. ropicalis</i> and one isolate of <i>C. glabrata</i> . When the biofilm activity and rate of acid production (ar.0228, pc.005) and thigmotropism (ar.0232, pc.004), but not cell surfaces were explored, a significant positive correlation was observed between the biofilm activity and rate of acid production (ar.0228, pc.005) and thigmotropism (ar.0232, pc.0401), but not cell surface bardrophobicity under 2028, pc.065) and thigmotropism (ar.0232, pc.0401), but not cell surface hydrophobicity (tri-0332, pc.0403). Taken together our in vitro data imply that candidal biofilm formation on aerylic surfaces is a complex phenomenon regulated by a multiplicity of factors operating intra-orally.