

IL-6 -174 Genotype Associated with A.actinomycetemcomitans in Indians

Luigi Nibali, Isobel Madden, Fernando Franch Chillida, Lisa Heitz-Mayfield,

Peter M Brett, Nikos Donos

▶ To cite this version:

Luigi Nibali, Isobel Madden, Fernando Franch Chillida, Lisa Heitz-Mayfield, Peter M Brett, et al.. IL-6 -174 Genotype Associated with A.actinomycetemcomitans in Indians. Oral Diseases, 2010, 17 (2), pp.232. 10.1111/j.1601-0825.2010.01731.x . hal-00599893

HAL Id: hal-00599893 https://hal.science/hal-00599893v1

Submitted on 11 Jun 2011

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers. L'archive ouverte pluridisciplinaire **HAL**, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d'enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.

ORAL DISEASES

IL-6 -174 Genotype Associated with A.actinomycetemcomitans in Indians

Journal:	Oral Diseases				
Manuscript ID:	ODI-04-10-OM-1648.R1				
Manuscript Type:	Original Manuscript				
Date Submitted by the Author:	18-May-2010				
Complete List of Authors:	Nibali, Luigi; UCL Eastman Dental Institute, Periodontology Madden, Isobel; 2 UHI School of Oral Health Science, Centre for Health Science Franch Chillida, Fernando; UCL Eastman Dental Institute, Periodontology Heitz-Mayfield, Lisa; The University of Western Australia, Centre for Rural and Remote Oral Health Brett, Peter; UCL Eastman Dental Institute, Periodontology Donos, Nikos; UCL Eastman Dental Institute, Periodontology				
Keywords:	Genetics, Periodontitis, Interleukin 6, Bacteria, A.actinomycetemcomitans				



IL-6 -174 Genotype Associated with A.actinomycetemcomitans in Indians

Running title: IL6 genotypes and A.actinomyctemcomitans

Nibali L.¹, Madden I.^{1, 2}, Franch-Chillida F.¹, Heitz-Mayfield L.J.A.³, Brett P.M.¹, Donos N.¹

¹ Periodontology Unit and Division of Clinical Research, Eastman Dental Hospital, University College London (UCL), London, UK.

² UHI School of Oral Health Science, Centre for Health Science, Inverness, Scotland

³ The University of Western Australia. Centre for Rural and Remote Oral Health. Crawley, Western Australia, Australia

Running title: IL-6 polymorphisms and periodontal bacteria

Keywords: Periodontitis, Interleukin-6, Genetic, Bacteria, A. actinomycetemcomitans

Date of submission: 22/04/2010

Corresponding author: Luigi Nibali Periodontology Unit Eastman Dental Institute 256 Gray's Inn Road London WC1X 8LD Tel:00442079151086 Fax 00442079151137 l.nibali@eastman.ucl.ac.uk

ABSTRACT

AIM: Genetic factors have recently been associated with presence of *Aggregatibacter actinomycetemcomitans* subgingivally in populations living in industrialised countries. The aim of this study was to analyze associations between Interleukin-6 (*IL6*) single nucleotide polymorphisms and presence and levels of *A.actinomycetemcomitans* and other subgingival microbes in a rural Indian population.

SUBJECTS AND METHODS: 251 individuals from a rural village in India with a periodontal phenotype ranging from healthy to severe periodontitis were included. Checkerboard DNA-DNA analysis was performed to detect 40 periodontal taxa in subgingival plaque samples. Genomic DNA was extracted to genotype five polymorphisms in the *IL6* promoter region.

RESULTS: The *IL6* -174 GG genotype was associated with high (above median) counts of *A.actinomycetemcomitans* (both in all subjects and in periodontally healthy only) and with presence and counts of *Capnocytophaga sputigena*. Differences in detection of several other bacteria were noted between periodontitis and healthy subjects.

factors on the s. CONCLUSION: These findings support the influence of genetic factors on the subgingival microbiota.

INTRODUCTION

The oral cavity may contain as many as 19,000 bacterial phylotypes (Kejiser et al., 2008). We have recently introduced the concept of periodontal infectogenomics, which studies the effect host genetic factors have on the composition of the subgingival microbiota of each individual. This is based on the assumption that genetic factors in the host seem to play a major role in deciding which bacteria (commensal and pathogenic) are able to colonise the host (Nibali et al., 2009a).

Most studies on subgingival microbes with a putative influence on periodontally-relevant immune responses have been carried out in industrialised communities with routine access to dental care (Hardie et al., 1975; Papapanou et al., 1993; Umeda et al., 1998). Analysis of rural populations not subjected to dental care allows the study of possible effects of host genetic factors in the composition of the subgingival microbiota in almost pristine conditions, undisturbed by routine tooth scaling and use of antibiotics.

We have recently shown an association between *IL6* genetic factors and clinical status in non smokers living in a rural Indian village (Franch et al., 2010). The main aim of this analysis was to study the association between *IL6* genetic variants and the presence and levels (counts) of *A.actinomycetemcomitans* in subgingival plaque samples from the same population. Secondary aims were to explore the association between *IL6* genetic factors and the detection and counts of other bacteria, and the association between periodontal status and the detection and levels of other bacteria.

MATERIALS AND METHODS

Subject Selection

The population reported here has also been described elsewhere (Franch et al., 2010). The study took place in the village of Dokur, Andhra Pradesh, where the Institute for Rural Health Studies (IRHS), a local Non-Governmental Organization (NGO), runs a rural clinic. All adult inhabitants between the ages of 18 and 70 were invited to attend a screening visit by means of local advertising within the village. A group of 500 villagers attended and were initially screened based on the following exclusion criteria: i) presence of serious systemic conditions (malaria, tuberculosis or cardiovascular disease), ii) presence of infectious or parasitic diseases, iii) pregnancy or breastfeeding, iv) antibiotic or anti-inflammatory medications within the last 3 months, v) fewer than 16 standing teeth.

Two hundred and fifty one suitable subjects (10% of the total village population) were identified and included in the study.

Ethical Considerations

The study was conducted in full accordance with ethics principles, including the World Medical Association Declaration of Helsinki (version 2002). Ethical approval for the study was obtained from the Medical Board of IRHS and conformed to the Indian Council of Medical Research Guidelines on Medical Ethics 2000 (Indian Council of Medical Research, Ansari Nager, New Delhi). All subjects were native to the village and voluntary informed consent to participation in the study was obtained from each by means of a written consent form, translated into Telugu and also read aloud to each subject by a fluent Telugu speaker. Demographic data and self-reported data on

education, social status, type of toothbrushing, alcohol consumption and smoking habit were recorded (Franch et al., 2010).

Clinical Data Collection

Full mouth periodontal examination of probing pocket depth (PPD) and clinical attachment level (CAL) was carried out for each subject with an EN-15 probe on 6 sites per tooth. The clinical examination was performed by one single operator (IM), calibrated to an exact kappa level of 0.85 for PPD. Patients were divided into 2 groups (periodontitis/ healthy) based on their periodontal status. Both definitions suggested by the European Workshop on Periodontitis (EWP) (Tonetti and Claffey, 2005) were used:

- Presence of proximal attachment loss of \geq 3mm in \geq 2 non-adjacent teeth (EWP1)
- Presence of proximal attachment loss of \geq 5mm in \geq 30% of teeth present (EWP2)

Microbiological analysis

Dental plaque samples were taken from the mesio-buccal aspects of the upper right and lower left first molar teeth, unless a deeper pocket had been found at another site, in which case this site was used in preference. Subgingival plaque was removed from each site isolated with cotton rolls, following removal of supragingival plaque, using sterile 13/14 double-ended Columbia curettes. The scraped mass was transferred from the curette to an Eppendorf tube containing 150µl of sterile TE buffer (10mM Tris HCl, 1.0mM EDTA, pH 7.6) and subsequently transferred to the University of Berne. The presence of 40 periodontal pathogens was evaluated using checkerboard DNA-DNA hybridisation technique (Socransky et al., 1994).

Genetic analysis



Samples of buccal mucosal cells were taken from all subjects using wire cytology brushes (Medical Wire and Equipment Company, UK) (Franch et al., 2010). Genomic DNA was extracted from these samples and blindly genotyped for polymorphisms at positions -174 (CCTTTAGCAT[C-G]GCAAGAC, rs 1800795), -572 (CAACAGCC[C-G]CTCACAG, rs 1800796), -1363 (CACTGTTTTATC[G-T]GATCTTG, rs 2069827) and -6106 (TCTCTACA[A-T]TAAGAAATAC) and -1480 (ACCGTCTCT[C-G]TGTTTAG) in the *IL6* gene by real time PCR as previously described (Nibali et al., 2008b).

Statistical Analysis

Comparisons of continuous and categorical data between groups (periodontitis and healthy) were analysed with ANOVA and Chi-square test, respectively. The alpha value was set at 0.05.

The approach to evaluate the association between genetic factors, clinical diagnosis and microbiological outcomes was performed using the SPSS 12.0 package. The main outcome was association between *IL6* -174 GG genotype and the presence (at detection threshold 1 x10⁴) and levels (total counts) of *A.actinomycetemcomitans*. An exploratory analysis was performed to investigate associations between all studied *IL6* genetic variables (and clinical diagnosis) and other taxa and bacterial complexes. The α value for this exploratory analysis was lowered to 0.01 in order to adjust for multiple testing. Microbiological data were available from two sites for all subjects. Data on detection or not of all 40 studied taxa are reported. For analysis of bacterial counts, negative (no detection) reactions were given an arbitrary value of 0.99 x10⁴, just below detection limit. The mean individual count for each species was obtained as the averaged between the 2 sites in each patient. A priori, median values for each bacterium were calculated and two clusters of patients for each bacterium were identified: high (above median) and low (below median) counts. Therefore, the microbiological outcomes were: i) detection of each bacterium (at 1 x10⁴ detection

Oral Diseases - Manuscript Copy

 value); ii) high or low levels (counts) of each bacterium (both categorical data) and iii) absolute counts of each bacterium (continuous data). Subgroup analyses were performed in subjects divided by their periodontal status (healthy/periodontitis). Multiple logistic regression analysis adjusting for confounders (age, smoking, gender and type of toothbrushing) was performed to investigate these associations (categorical variables). Linear logistic regression adjusting for confounders (age, smoking, gender and type of toothbrushing) was used to detect differences in bacterial counts (continuous variables) between genotypes and by clinical diagnosis,

Deleted: Mann-Whitney test

Deleted: as previously described (continuous variables) (Socransky et al., 2000)

RESULTS

A total of 251 subjects took part in the study. Their demographic and clinical characteristics have been reported elsewhere (Franch et al., 2010) (table 1). Forty-two per cent and 19% of subjects were diagnosed with periodontitis, according to EWP1 and EWP2 definitions respectively (Tonetti and Claffey, 2005). Thirteen subjects resulted *IL6* -174 C homozygous, 46 CG and 185 GG, while 7 could not be scored. The distribution of genetic polymorphisms has been reported elsewhere and satisfied the chi-squared analysis for Hardy-Weinberg equilibrium on all subjects (Franch et al., 2010).

The detection of the 40 studied taxa ranged from 98% of subjects for *Prevotella intermedia* to 51% for *Capnocytophaga sputigena*. In particular, 93% of subjects were positive for *A. actinomycetemcomitans*, 93% for *Porphyromonas gingivalis*, 90% for *Tannerella forsythia*, 70% for *Treponema denticola* and 60% for *Campylobacter rectus*. The mean counts ranged from $11_2^2 \times 10^5$ for *Neisseria mucosa* to 0.2×10^5 for *Propionibacterium acnes*. Average counts for *A. actinomycetemcomitans*, *P. gingivalis* and *T. forsythia* for all subjects were respectively $5_2^1 \times 10^5$, 12.0×10^5 and 6.4×10^5 .

Deleted: periodontally healthy

Association between IL6 genetic factors and bacterial detection

IL6 -174 GG genotype subjects exhibited high (above median) levels of *A.actinomycetemcomitans* compared to CC and CG individuals, irrespective of clinical diagnosis (adjusted p=0.043, O.R. = 1.89, 95% C.I. = 1.02- 3.50) (figure 1). This association was confirmed among the subgroup of <u>non-periodontitis</u>, individuals (according to EWP2) (adjusted p=0.037, O.R. = 2.11, 95% C.I. = 1.0-4.2).

Among other bacteria, *IL6* -174 GG genotype subjects showed increased detection, and high (above median) levels of *C.sputigena* (adjusted p= 0.003, O.R. = 2.60, 95% C.I. = 1.38- 4.90 and p=0.009, O.R. = 2.29, 95% C.I. = 1.23- 4.26 respectively). No statistically significant differences in percentage of bacterial complexes (Socransky *et al.*, 1998) were found according to IL6 -174 genotypes (figure 2).

Associations between clinical diagnosis and bacterial detection

Figure 3 shows the frequency of detection of the 40 studied bacteria in periodontitis and healthy individuals according to EWP1 (Tonetti and Claffey, 2005). Statistically significant associations were found between diagnosis of periodontitis (EWP1) and presence of *N.mucosa* (p=0.008) and *Capnocytophaga ochracea* (p=0.001), low (below-median) counts of *N.mucosa* (p<0.001), and high (above-median) counts of *Parvimonas micra* (previously *Peptostreptococcus micros*) (p=0.002). Periodontitis patients also exhibited lower absolute counts of *N.mucosa* (p=0.003).

EWP2 diagnosis of periodontitis was associated with absence (no detection) of *C.ochracea* (p=0.005) and *S.noxia* (p<0.001) and higher absolute counts of *P.micra* (p=0.025). When bacteria were grouped in complexes, periodontal status (EWP1) was associated with percentage of bacteria undefined in complexes ('others') (p=0.012).

Deleted: 8

Deleted:), *C.ochracea* (p<0.001), and *Selenomonas noxia* (p<0.001)

Deleted: os

Deleted: <0.001

Deleted: <0.001 and p=0.001 for EWP1 and EWP2 respectively

Deleted: EWP2 definition was also associated with percentage of green complex bacteria (p=0.002).

Deleted: subgingival

DISCUSSION

We have previously shown an association between *IL6* -174 genotypes and presence of periodontitis in non-smokers living in a rural village in India (Franch et al., 2010). The present study describes associations between *IL6* -174 GG genotype and presence of *A.actinomycetemcomitans* in periodontal pockets in these subjects (with a periodontal phenotype ranging from healthy to severe periodontitis), confirming association previously observed in subjects living in the United Kingdom (Nibali et al., 2007, Nibali et al., 2008a). Among these Indian villagers living in Andra Pradesh, *A.actinomycetemcomitans* was almost ubiquitous subgingivally, as it was harboured by 93% of participating individuals. However, *IL6* -174 GG individuals had higher counts of this bacterium.

The *IL6* -174 GG has been associated with presence of periodontitis (Trevilatto et al., 2003, Nibali et al., 2009b). The study reported here confirms the previously reported association between *IL6* genotypes and *A.actinomycetemcomitans* (Nibali et al., 2007, Nibali et al., 2008a), despite being conducted on a different population of different ethnicity and socio-economic status and not subjected to routine dental care. Furthermore, the microbiological analysis consisted of a checkerboard DNA-DNA hybridisation analysis (Socransky et al., 1994), as opposed to previously reported culture (Nibali et al., 2007) and PCR (Nibali et al., 2008a) analyses. This association was seen in all subjects, independent from clinical diagnosis. In addition to this, this paper for the first time shows the association between IL6 genotypes and *A.actinomycetemcomitans* also in non-periodontitis, individuals (according to EWP2 definition). As previously hypothesized, this association may be due to the increased local inflammatory response of -174 GG individuals (Fishman et al., 1998, Bennermo et al., 2004), which favours the overgrowth of components of the microbiota that grow well in inflamed areas, such as *A.actinomycetemcomitans*. This study brings preliminary evidence that other bacteria belonging to the green complex, in particular *C.sputigena*,

Deleted: periodontally healthy

Oral Diseases - Manuscript Copy

Deleted: subgingival

may also be associated with increased counts in <u>periodontal pockets of *IL6*-174 GG individuals. In</u> other terms, the effect of genetic factors may not be limited to the presence or levels of one or two potentially pathogenic bacteria, but may extend to bacterial complexes or in general to the whole composition of the subgingival biofilm. This is in agreement with the concept of infectogenomics, suggesting that genetic factors in the host seem to play a major role in deciding which bacteria (commensal and pathogenic) are able to colonise the host (Kellam and Weiss, 2006, Nibali et al.,

2009a).

The microbiological analysis of the current subject group showed a population presenting with a complex oral flora, with a breakdown in bacterial complexes quite different from that observed in industrialized populations (Haffajee et al., 1998). Individuals diagnosed with periodontitis had a tendency towards higher percentage of green complex bacteria, and lower percentage of bacteria not defined in complexes ('others'). Overall, there was a predominance of bacteria belonging to the orange complex, which have been associated with established periodontitis in industrialised populations, while no significant differences in percentage of orange or red complex bacteria were noted between periodontitis and non-periodontitis individuals. This was consistent with studies in other untreated populations, which have shown ubiquitous presence of certain bacterial species and a tendency towards a stable, mixed microflora (Timmerman et al., 1998, Dowsett et al., 2002, Papapanou et al., 2002, Colombo et al., 2002). N.mucosa, a Gram negative commensal bacterium, was the bacterium found in highest counts, followed by periodontopathogenic bacteria *P. gingivalis*, *T.forsythia* and *A.actinomycetemcomitans*. This shows a considerable shift from the bacterial profile shown in a study with checkerboard DNA-DNA hybridization on a large United States population (Haffajee et al., 2006). The difference in prevalence and counts of A. actinomycetem comitans between the study described here and the one by Haffajee and coworkers is particularly striking. It needs to be highlighted that only two of the deepest pockets per subject were sampled in the present

study. Therefore, the average bacterial counts in these cases in not representative of the whole subgingival microbiota. However, even when the sampled site was used as unit for analysis, no differences were detected between bacterial complexes between deep (>6mm) and shallow (<4mm) pockets (data not presented).

Among the drawbacks of this study, we have to acknowledge the limitation of having only two plaque samples sites per patient, the risk for false positive results associated with DNA-DNA checkerboard analysis (van Steenbergen et al., 1996) and the difficulty of defining periodontitis in this population (Preshaw 2009, Franch et al. 2010). The results reported here show an association between *IL6* -174 polymorphism and *A.actinomycetemcomitans* in subjects living in a rural Indian village, bringing further evidence, consistent with previous studies, of the importance of infectogenomics in periodontitis (Nibali et al., 2009a). Further studies should analyze the possible effect of these and other genetic factors (in both periodontitis and healthy populations) on a wider range of components of the oral microbiota, on different serotypes and clones of *A.actinomycetemcomitans*, and should investigate possible functional mechanisms for the observed associations.

ACKNOWLEDGEMENTS

This work was undertaken at UCLH/UCL who received a proportion of funding from the Department of Health's NIHR Biomedical Research Centres funding scheme. This study was also supported by the Periodontal Research Fund of the Eastman Dental Institute. The help of Professor Gareth Griffiths, Professor Hubert Newman, Dr Bidinger and the Institute for Rural Health Studies, Hyderabad, Andhra Pradesh, are gratefully acknowledged.

Deleted: (which might to some extent affect the overall bacterial counts)

REFERENCES

Bennermo M, Held C, Stemme S, Ericsson CG, Silveira A, Green F et al. (2004). Genetic predisposition of the interleukin-6 response to inflammation: Implications for a variety of major diseases? Clinical Chemistry **50**: 2136-2140.

Fishman D, Faulds G, Jeffery R, Mohammed-Ali V, Humphries S, Woo P (1998). An interleukin-6 promoter polymorphism that influences gene transcription, plasma IL-6 levels and is associated with systemic-onset juvenile chronic arthritis. European Cytokine Network **9**: 364.

Kellam P, Weiss RA (2006). Infectogenomics: Insights from the host genome into infectious diseases. Cell **124**: 695-697.

Keijser BJF, Zaura E, Huse SM, van der Vossen JMBM, Schuren FHJ, Montijn RC et al. (2008). Pyrosequencing analysis of the Oral Microflora of healthy adults. Journal of Dental Research **87**: 1016-1020.

Colombo APV, Teles RP, Torres MC, Souto R, Rosalem W, Mendes MCS et al. (2002). Subgingival microbiota of Brazilian subjects with untreated chronic periodontitis. Journal of Periodontology **73**: 360-369.

Dowsett SA, Kowolik MJ, Archila LA, Eckert GJ, LeBlanc DJ (2002). Subgingival microbiota of indigenous Indians of Central America. Journal of Clinical Periodontology **29**:159-167.

Franch Chillida F, Nibali L, Madden I, Donos N, Brett PM (2009). Association between Interleukin-6 polymorphisms and periodontitis in a rural Indian population. Journal of Clinical Periodontology **37**: 137-144.

Hardie JM, Bowden GH (1975). Bacterial-Flora of Dental Plaque. British Medical Bulletin **31**:131-136.

Haffajee AD, Cugini MA, Tanner A, Pollack RP, Smith C, Kent RL et al. (1998). Subgingival microbiota in healthy, well-maintained elder and periodontitis subjects. Journal of Clinical Periodontology **25**:346-353.

Haffajee AD, Teles RP, Socransky SS (2006). Association of Eubacterium nodatum and Treponema denticola with human periodontitis lesions. Oral Microbiology and Immunology **21**:269-282.

Nibali L, Ready DR, Parkar M, Brett PM, Wilson M, Tonetti MS et al. (2007). Gene polymorphisms and the prevalence of key periodontal pathogens. Journal of Dental Research **86**:416-420.

Nibali L, Tonetti MS, Ready D, Parkar M, Brett PM, Donos N et al. (2008a). Interleukin-6 polymorphisms are associated with pathogenic bacteria in subjects with periodontitis. Journal of Periodontology **79**:677-683.

Nibali L, Griffiths GS, Donos N, Parkar M, D'Aiuto F, Tonetti MS et al. (2008b). Association between interleukin-6 promoter haplotypes and aggressive periodontitis. Journal of Clinical Periodontology **35**:193-198.

- Nibali L, Donos N, Henderson B (2009a). Periodontal infectogenomics. Journal of Medical Microbiology **58**:1269-1274.
- Nibali L, D'Aiuto F, Donos N, Griffiths GS, Parkar M, Tonetti MS et al. (2009b). Association between periodontitis and common variants in the promoter of the interleukin-6 gene. Cytokine **45**:50-54.
- Papapanou PN, Sellen A, Wennstrom JL, Dahlen G (1993). An Analysis of the Subgingival Microflora in Randomly Selected Subjects. Oral Microbiology and Immunology 8:24-29.
- Papapanou PN, Teanpaisan R, Obiechina NS, Pithpornchaiyakul W, Pongpaisal S, Pisuithanakan S et al. (2002). Periodontal microbiota and clinical periodontal status in a rural sample in southern Thailand. European Journal of Oral Sciences **110**:345-352.
- Preshaw PM (2009). Definitions of periodontal disease in research. Journal of Clinical
 Periodontology 36:1-2.
 - Socransky S, Smith AC, Martin L, Paster BJ, Dewhurst FE, Levin AE (1994). Checkerboard DNA-DNA hybridisation. Biotechniques 17: 788 792.
 - Socransky SS, Haffajee AD, Cugini MA, Smith C, Kent RL (1998). Microbial complexes in subgingival plaque. Journal of Clinical Periodontology **25**:134-144.
 - Socransky SS, Haffajee AD, Smith C, Duff GW (2000). Microbiological parameters associated with IL-1 gene polymorphisms in periodontitis patients. Journal of Clinical Periodontology **27**:810-818.
 - Timmerman MF, Van der Weijden GA, Abbas F, Arief EM, Armand S, Winkel EG et al. (2000). Untreated periodontal disease in Indonesian adolescents - Longitudinal clinical data and prospective clinical and microbiological risk assessment. Journal of Clinical Periodontology **27**:932-942.
 - Tonetti MS, Claffey N. European Workshop in Periodontology group C. (2003) Advances in the progression of periodontitis and proposal of definitions of a periodontitis case and disease progression for use in risk factor research. Group C consensus report of the 5th European Workshop in Periodontology. Journal of Clinical Periodontology **32** Suppl.: 210-3
 - vanSteenbergen TJM, Timmerman MF, Mikx FHM, deQuincey G, vanderWeijden GA, vanderVelden U et al. (1996). Discrepancy between culture and DNA probe analysis for the detection of periodontal bacteria. Journal of Clinical Periodontology **23**:955-959.
 - Umeda M, Chen C, Bakker I, Contreras A, Morrison JL, Slots J (1998). Risk indicators for harboring periodontal pathogens. Journal of Periodontology **69**:1111-1118.

FIGURE LEGENDS

Figure 1. Representation of number of subjects with A.actinomycetecomitans above or below median according to their IL-6 -174 genotypes. GG vs CG/CC p=0.043, O.R.= 1.89, 95% C.I.=1.02-3.50.

Figure 2. Pie charts with distribution of bacterial complexes between IL6 -174 GG and CG/CC individuals. % DNA probe counts for each bacterium was determined at each site, and averaged per patient. The % of the DNA probe count for each species in each complex (according to Socransky et al., 1998) was summed and the proportions of each complex were calculated.

Figure 3. Counts of 40 studied taxa divided in complexes in healthy and periodontitis subjects (according to EWP1 definition).

* *P.micra*: p<0.01 for differences in high (above median) counts between healthy and periodontitis

** C.ochracea: p<0.01 for differences in detection and high (above median) counts between healthy and periodontitis

*** N.mucosa: p<0.01 for differences in detection, high (above median) counts and absolute counts between healthy and periodontitis.

		All subjects (n=251)	1	EWP1	EWP2		
			Periodontitis (n=105)	Non- periodontitis (n=146)	Periodontitis (n=47)	Non-periodontitis (n=204)	
Age			45.8 ± 10.5	37.5± 11.1	49.6 ± 10.3	39.0±10.9	
Gender (female)		162 (64.5%)	62 (59.0%)	100 (68.5%)	28 (59.6%)	134 (65.7%)	
Smoking	Current	44 (17.5%)	26 (24.8%)	18 (12.3%)	33(70.2%) 14 (29.8%)	174(85.3%)	
	Never	207 (82.5%)	79 (75.2%)	128(87.7%)		30(14.7%)	
Clinical data	Number of sites PPD 4- 6mm	16.1 ± 22.1	38.5 ± 17.4	0.1 ± 0.4	45.5±15.9	8.7±15.6	
	Number of sites PPD >6mm	3.3 ± 8.1	7.9 ± 11.1	0.0 ± 0.2	15.1±13.1	0.6±1.8	
	Number of sites CAL >4mm	16.6 ± 22.1	39.5±16.3	0.1 ± 0.6	47.5±15.5	9.5±16.6	
	Number of sites CAL	6.1±14.0	14.5±18.5	0.0± 0.4	27.5±21.1	1.1±2.7	

Table 1. Demographic and clinical characteristics of the participants, as divided by their periodontal status.

Deleted: * * <i>S.noxia</i> : p<0.01 for differences in total counts between
healthy and periodontitis
Deleted: *

Deleted: os

Page 15 of 17





COUNTS 10 ⁵	0 2	4	6	8	10	12	14	16
A. gerencseriae	-	1	1			AC	TINOMYCES	
A. israelii	1							
A. naeslundii 1								
A. naeslundii 2								
A. odontolyticus							FORFLE	
V. parvula								
S. gordonii								
S. internedius							YELLOW	
S. mitis		>						
S. oralis								
S. sanguinis								
A. actinomycetemcomitans			>					
C. gingivalis							GREEN	
C ochrace a]** 🔣							
C. sputigen a								
E corrodens								
C gracilis								
C. rectus								
C. showae	1						ORANGE	
E nodatum								HEALT
F. nucleatum ss nucleatum								(n=146
F. nucleatum ss polymorphum								
F. nucleatum ss vincentii	-							
F. periodonticum		_						
P. micros]* 🦰							
P. intermedia								(n=105)
P. nigrescens								(11-105)
S. constellatus								
T. forsythia								
P. gingivalis							RED	
T. denticola								
E. saburreum								
G. morbillorum							OTHERS	
L. buccalis								
N. mucosa	***							
P. acnes								
P. melanin ogenica								
S. anginosus								
S. noxia		>						

