Research Article

Clonal Relationship within Two Oral Actinomyces Populations Collected from Plaque of Periodontitis Patients

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A B S T R A C T

Actinomyces naeslundii and A. oris are dental plaque formers involved in the pathogenesis of periodontitis. The aim of the study was to investigate the clonal relationship within two oral Actinomyces populations collected from plaque of patients with chronic periodontitis. The 223 clinical strains of A. naeslundii and A. oris were isolated from biofilm samples collected supra and subgingivally from teeth with shallow (probing depth (PD) = 3-4 mm), deep (PD = 5-6 mm) and very deep (PD ≥7 mm) pockets from 20 chronic periodontitis patients. All strains were submitted to repetitive sequence-based PCR typing using DiversiLab (BioMerieux, Marcy l’Étoile, France). Seven patients harboured only unrelated (>95% similarity) multiple isolates, while 13 harboured both similar (>95% similarity) and unrelated isolates at different sites. Identical (>98% similarity) strains were found to be present in the subgingival shallow depths more often than in the other subgingival depths. The number of clones in individual patients varied from 2 to 17 different rep-PCR genotypes. The clonal relationship within the oral populations of A. naeslundii and A. oris in an individual was unpredictable, ranging from the presence of multiple genotypes with no clonal similarity to only two different clones supra or subgingivally at different sites.

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Introduction

Actinomyces spp. are gram-positive, non-motile, non-spore-forming, and pleomorphic rods [1]. Currently, Actinomyces spp. comprise 42 species, 20 of which are considered relevant to human medicine [2-4]. They are commonly part of the flora of the respiratory tract, oral cavity, and other habitats like the genito-urinary system and the skin [4, 5]. They are also associated with infections such as cervicofacial and hepatic actinomycosis, cerebral or oral abscesses, urinary tract infections, caries, and periodontitis [1, 6-11]. Actinomyces spp. act as early colonisers in the process of plaque maturation [11, 12]. They adhere to the tooth surface and provide a substrate for the adherence of other plaque microorganisms [13, 14]. Together with other bacteria, they form supra and subgingival polymicrobial biofilms, which are involved in the pathogenesis of different forms of periodontitis [12, 15-21]. Recent studies have shown that Actinomyces naeslundii can induce horizontal alveolar bone loss and can protect many bacterial species from hydrogen peroxide [22, 23]. Vielkind et al. determined a massive presence of A. naeslundii/oris/johnsonii that outnumbered all the other Actinomyces species in almost all the supra and subgingival/sulcular samples from both periodontitis patients and healthy subjects [24]. These findings support the fact that these species are probably the most important plaque formers among the members of the Actinomyces genus. Most of the previous studies have reported only the composition of supra and subgingival biofilms up to the species level, while the heterogeneity of the individual strains remains unexplored. The emergence of the automated repetitive sequenced-based PCR bacterial typing system, DiversiLab offers the possibility of such studies. Recent studies of the clonal diversity of Streptococcus mutans of household family members showed that DiversiLab is used to investigate the genetic diversity [25]. DiversiLab is known for sim...
A. oris populations isolated from different depths of periodontal pockets in patients with chronic periodontitis.

Materials and Methods

I Bacterial Strains and Culture Conditions

A total of 223 Actinomyces strains (A. oris (44) and A. naeslundii (179)) were used in this study. They were isolated from the supra and subgingival biofilms of 20 periodontitis patients. Subgingivally, the samples were collected from teeth with shallow pockets (probing depth (PD) of 3-4 mm), deep pockets (PD of 5-6 mm), and very deep pockets (PD ≥ 7 mm). Supragingivally, the samples were collected from the sites corresponding to the previously mentioned pocket depths. The strains were frozen at -80°C in skimmed milk. Prior to use, the bacteria were cultured for 4 days on Colombia Blood Agar Base (Oxoid Microbiology Products, Thermo Fisher Scientific, United Kingdom) at 37°C in an anaerobic chamber (Whitley MG1000 anaerobic workstation, Meintrup DWS Laborgeräte, GmbH, Germany). All the strains were identified using MALDI-TOF-MS (Autoflex II, Bruker Daltonics, Billerica, MA) and by the sequence analysis of the 16S ribosomal RNA (rRNA) gene [35].

II rep-PCR

i DNA Extraction

DNA extraction from the cultivated strains was performed employing a MagNA Pure 96 System (Roche Molecular Systems, Rotkreuz ZG, Switzerland) according to the manufacturer’s instructions.

ii Automated rep-PCR DNA Fingerprinting

rep-PCR typing was performed using DiversiLab 3.6.0.39 (BioMérieux, Marcy l’Étoile, France), an automated repetitive sequence-based PCR bacterial typing system. This system allows the amplification of many different-sized fragments (amplicons) representing the DNA within the non-coding, repetitive sequences in the genome. rep-PCR primers bind to many specific repetitive genome sequences and multiple fragments of various lengths are amplified. These fragments can be separated by size and charge using microfluidics electrophoresis. A unique rep-PCR fingerprint profile is created containing multiple bands of varying sizes and intensities. Inside the LabChip® (DiversiLab, BioMérieux), the DNA fragments are allowed to bind with a dye, and a graph of fluorescence intensity over time is created. This graph translates into the fingerprint pattern of the sample. First, the purified DNA was quantified with a Nanodrop 1000 spectrophotometer (Thermo Fisher Scientific). The rep-PCR amplification was performed with the designated DiversiLab fingerprinting kit and sizing was performed using a microfluid LabChip. The chip was placed into a 2100 Bioanalyzer (Agilent Technologies, Software Version 3.4.) and the electrophoresis was performed automatically.

iii Statistical Analysis

Electrophoresis results were uploaded to the DiversiLab Analysis with the help of the BioMérieux Customer Help. The data were analysed using the Pearson’s Correlation test. A colour-code emphasises and assists in reading and interpreting the results of the electrophoresis. Furthermore, a dendogram (Figure 1) is provided, which shows the relationship between the isolates of a patient from different pocket depths. Based on the work of Paul G. Higgins, we decided on the following similarity rules [36]:

i. Strains that exhibited <95% similarity were classified as unrelated.

ii. Strains that exhibited a similarity between 95%-98% were classified as closely related isolates.

iii. Strains that exhibited >98% similarity were classified as identical isolates.

Figure 1: Dendrogram of Actinomyces naeslundii of patient 16 and electropherogram of the rep-PCR. The strains numbered 1 to 13 showed a clonal similarity >98%, (colour-code: dark red) including supragingival (P16B4, P16A5, P16C4, P16A2, P16C2, P16A11, P16B5) and subgingival sites (P16D11, P16D2, P16D4, P16F7, P16D10, P16F8,). Strains numbered 14 to 18 (P16D6, P16F4, P16B9, P16B10, P16C1) did not show any clonal relationships (colour-code: orange (<98%), blue (<90 %), yellow (<80%), grey (<70%).
The clonal diversity of *Actinomyces* was analysed for each patient individually. All the comparisons were performed for strains isolated from different sites of a single patient. To assess the reliability of the DiversiLab system, we also compared specimens from different patients harbouring the same species.

**Results**

The distribution of *Actinomyces* strains in the supra and subgingival plaques of periodontitis patients at different pocket depths is shown in (Table 1). As patient no. 14 did not harbour any *A. naeslundii* or *A. oris* strains, pertaining data is not included in (Table 1). Patient no. 6 did not harbour any *A. naeslundii* strains. *A. naeslundii* and *A. oris* strains were found either supra or subgingivally or only supragingivally or on both sites. The clonal similarity between all the *A. naeslundii* and *A. oris* strains is shown in (Table 2). Multiple colonisation patterns were found. Seven patients (1, 2, 4, 5, 8, 12, and 13) were individually colonised with only unrelated isolates (similarity <98%), while 13 patients harboured both similar and unrelated isolates at different sites. Figure 1 shows the clonal relationship within the *A. naeslundii* population isolated from the patient 16. This patient harboured multiple clones but one of them was predominant and found at many sites. Figure 2 shows the clonal relationship within the population of *A. naeslundii* and *A. oris* isolated from the patient 13. This patient harboured only unrelated multiple clones of both species.

Table 1: Distribution of *A. naeslundii* and *A. oris* strains the in supra and subgingival biofilm.

| Patient | Species     | Total strains | Supragingival sites corresponding to the following pocket depths | Subgingival |
|---------|-------------|---------------|-----------------------------------------------------------------|-------------|
|         |             |               | 3-4 mm (A) | 5-6 mm (B) | ≥7 mm (C) | 3-4 mm (D) | 5-6 mm (E) | ≥7 mm (F) |
| 1       | *A. naeslundii* | 4             | 2         | 0         | 0         | 0         | 2         | 0         |
|         | *A. oris*   | 2             | 2         | 0         | 0         | 0         | 0         | 0         |
| 2       | *A. naeslundii* | 5             | 3         | 0         | 0         | 1         | 1         | 0         |
|         | *A. oris*   | 0             | 0         | 0         | 0         | 0         | 0         | 0         |
| 3       | *A. naeslundii* | 17            | 4         | 5         | 1         | 2         | 1         | 4         |
|         | *A. oris*   | 0             | 0         | 0         | 0         | 0         | 0         | 0         |
| 4       | *A. naeslundii* | 7             | 1         | 2         | 3         | 1         | 0         | 0         |
|         | *A. oris*   | 0             | 0         | 0         | 0         | 0         | 0         | 0         |
| 5       | *A. naeslundii* | 5             | 1         | 2         | 2         | 0         | 0         | 0         |
|         | *A. oris*   | 1             | 1         | 0         | 0         | 0         | 0         | 0         |
| 6       | *A. naeslundii* | 0             | 0         | 0         | 0         | 0         | 0         | 0         |
|         | *A. oris*   | 11            | 2         | 4         | 0         | 0         | 2         | 3         |
| 7       | *A. naeslundii* | 10            | 2         | 2         | 1         | 3         | 2         | 0         |
|         | *A. oris*   | 3             | 0         | 0         | 2         | 0         | 0         | 1         |
| 8       | *A. naeslundii* | 3             | 0         | 1         | 0         | 2         | 0         | 0         |
|         | *A. oris*   | 0             | 0         | 0         | 0         | 0         | 0         | 0         |
| 9       | *A. naeslundii* | 9             | 4         | 1         | 3         | 1         | 0         | 0         |
|         | *A. oris*   | 1             | 0         | 0         | 0         | 0         | 0         | 0         |
| 10      | *A. naeslundii* | 10            | 3         | 2         | 2         | 3         | 0         | 0         |
|         | *A. oris*   | 0             | 0         | 0         | 0         | 0         | 0         | 0         |
| 11      | *A. naeslundii* | 11            | 3         | 4         | 2         | 1         | 1         | 0         |
|         | *A. oris*   | 0             | 0         | 0         | 0         | 0         | 0         | 0         |
| 12      | *A. naeslundii* | 10            | 1         | 1         | 4         | 1         | 1         | 2         |
|         | *A. oris*   | 1             | 0         | 1         | 0         | 0         | 0         | 0         |
| 13      | *A. naeslundii* | 14            | 0         | 3         | 5         | 2         | 4         | 0         |
|         | *A. oris*   | 3             | 0         | 0         | 2         | 1         | 0         | 0         |
| 14      | *A. naeslundii* | 9             | 3         | 1         | 3         | 0         | 1         | 1         |
|         | *A. oris*   | 6             | 2         | 2         | 0         | 2         | 0         | 0         |
Table 2: Clonal diversity of *Actinomyces* strains in the supragingival and subgingival biofilms of patients.

| Patient | Species | Clones | Genotypes of A. naeslundii/A. oris |
|---------|---------|--------|-----------------------------------|
|         |         |        | Supragingival | Subgingival | Supra and subgingival |
| 1,2,4,5 | A. naeslundii | - | - | - | - |
|         | A. oris | - | - | - | - |
| 3,11    | A. naeslundii | - | - | - | - |
|         | A. oris | - | - | - | - |
| 6       | A. naeslundii | - | - | - | - |
|         | A. oris | - | - | - | - |
| 7       | A. naeslundii | - | - | - | - |
|         | A. oris | - | - | - | - |
| 9       | A. naeslundii | - | - | - | - |
|         | A. oris | - | - | - | - |
| 10      | A. naeslundii | - | - | - | - |
|         | A. oris | - | - | - | - |
| 8,12,13 | A. naeslundii | - | - | - | - |
|         | A. oris | - | - | - | - |
| 15      | A. naeslundii | - | - | - | - |
|         | A. oris | - | - | - | - |
| 16,17,19| A. naeslundii | - | - | - | - |
|         | A. oris | - | - | - | - |
| 18      | A. naeslundii | - | - | - | - |
|         | A. oris | - | - | - | - |
| 20      | A. naeslundii | - | - | - | - |
|         | A. oris | - | - | - | - |

*"-" = unrelated strains with a similarity of < 95%

*"+" = identical isolates with a similarity of > 98%
Figure 2: Dendrogram of patient 13 and electropherogram of the rep-PCR. Strains 2 and 5 to 17 were identified as *Actinomyces naeslundii*; 1, 3, and 4 as *Actinomyces oris*. No strains showed any clonal relationship (colour-code: orange (<98%), blue (<90 %), yellow (<80%), grey (<70%). Strains 16 and 17 have a similarity of 97.8% (colour-code: red).

Figure 3: Dendrogram of *Actinomyces naeslundii* of patient 3 and the electropherogram of the rep-PCR. Strains 1 to 3 (P3B9, P3C2, P3B11) showed a clonal relationship just supragingivally. Strains 7 to 14 (P3A1, P3A2, P3A11, P3B3, P3B5, P3F15, P3F1, P3F3) showed similarity in supra and subgingival depths. Strains 4-6 (P3D6, P3B8/1, P3D3) and 15-17 (P3A4/2, P3F10, P3E9) showed no clonal relationship (colour-code: orange (<98%), blue (<90 %), yellow (<80%), grey (<70%).

Figure 3 shows the clonal relationship within the population of *Actinomyces naeslundii* strains isolated from patient No. 3. This patient harboured multiple similar clones, both supra and subgingivally. Similar clones within a population were found either supra or subgingivally or both. No difference between the colonisation patterns of the two species was found. Some patients harboured identical clones of either *A. oris* or *A. naeslundii* only supragingivally while others harboured both supra and subgingivally. In most of the patients, we observed that the supragingival clones were also present subgingivally. Interestingly, two patients were colonised with similar clones both, supra and subgingivally, but these clones were unrelated to each other. The similar clones found subgingivally were not always found at all the pocket depths. Identical strains occurred in subgingival shallow depths more often than in the other subgingival depths. The number of clones in one patient varied from 2 to 17 different rep-PCR genotypes. No similarity was found between the strains isolated from different patients.
Discussion

Oral microflora is very complex and exposed to several individual factors such as nutrition, hygiene, and habits [37, 38]. The composition changes during the different stages of life, influenced by medication, health, and general constitution. 

Actinomyces spp. are early colonisers in the development of oral plaques and can occur in periodontal pockets [24]. During the plaque maturation, they contribute to the coaggregation with other bacteria [39]. A. naeslundii and A. oris possess fimbriae that adhere to the host surface and build up cell-to-cell interaction, especially with oral Streptococci [40]. Cluster analysis of supragingival biofilms has shown that Actinomyces spp. form an Actinomyces-complex in supragingival biofilms with a close relationship to the yellow complex bacteria, that mainly consists of Streptococci [19]. They occur both in healthy subjects and in patients with periodontal disease [41]. A high prevalence of Actinomyces is also shown in peri implantitis-sites [42]. In this study, only A. naeslundii and A. oris were examined since they always occur in oral biofilms and periodontal pockets in high concentrations in almost all the patients [43]. A. naeslundii decreases in quantity from shallow to deep pockets in the process of periodontitis [35]. The clonal relationship within the population of A. naeslundii and A. oris at different depths of periodontal pockets and supragingivally on the sites corresponding to these depths was performed using DiversiLab, an automated repetitive sequenced-based PCR bacterial typing system. In comparison to pulsed-field gel electrophoresis (PFGE), DiversiLab is less time-consuming and more convenient [44].

DiversiLab has shown more practicability in comparison with PFGE but results in more indistinguishable strains and is less discriminatory [28, 44-46]. It has been shown to overestimate the genetic relatedness in comparison with PFGE and to include more isolates to be part of a cluster when compared with that of AFLP (Amplified fragment length polymorphism). However, AFLP is more labour intensive [46, 47]. Overall, DiversiLab was a user-friendly and a very convenient typing method within the scope of this study. The software allows an easy comparison of numerous probes. Compared to AFLP and PFGE, less time has to be invested and the reproducibility is also good. The analysing programme (DiversiLab Software v3.4.4) provides three options for investigating similarities of various specimens: the modified Kullback-Leibler, the extended Jaccard, and the Pearson’s correlation. These options rely on the relative intensities of the sample pair at each data point. For this study, the Pearson’s correlation was chosen. This method emphasises peak intensities rather than peak presence or absence. This is helpful when there are many different variations in the fingerprint patterns. The comparison is shown in a matrix with percentage numbers that express the similarity of the fingerprint patterns. Thus, samples of every chip that were assessed by the system could be compared with each other. DiversiLab is able to generate genotypic profiles by using a nanogram scale of DNA and is able to compare data longitudinally and potentially between laboratories [48].

Given the complexity of the oral microflora, one might expect to find clonal diversity within the members of a species. The high prevalence of A. naeslundii and A. oris supra and subgingivally offered the possibility to study this hypothesis. The results were diverse as multiple colonisation patterns were observed. Some patients harboured multiple genotypes of A. naeslundii and/or A. oris. For instance, patient no. 13 (Figure 2) showed a high clonal diversity harbouring 17 different genotypes. Other patients showed a less striking diversity with only two genotypes in each: patient no. 16 showed 2 genotypes of A. naeslundii, while patient 20 showed 2 genotypes of A. oris. Other patients, although colonised with more than one clone, harbour a single clone also on different sites. Our results showed that identical clonal strains occur in subgingival shallow depths more often than in the other subgingival depths. Supragingivally, the majority of the strains were found in the sites corresponding to the pockets of depths 5-6mm. In our patients, a common distribution pattern of identical isolates was not detected. In some patients, identical isolates were found supragingivally while subgingivally, they were colonised with different clones. The distribution of identical isolates also showed a high variability. For example, one clone isolated thrice supragingivally in patient no. 3 was also isolated subgingivally, but together with other isolates which were not similar to the initial ones. However, in patient no. 7, we found the identical clone five times supragingivally and only once together with other different clones subgingivally.

Actinomyces spp. are facultative anaerobic bacteria, which can not only survive in the deep pockets, but also in oxygen-rich supragingival biofilms. This leads to a suggestion that clones may adapt to different environmental circumstances as well as possibly to different forms of therapy. The bacteria might possess various virulence factors and develop resistance mechanisms against antibiotic agents. Thus, further studies should include various clones when investigating these properties. The clonal relationship within the population of A. naeslundii and A. oris in an individual is variable and unpredictable. It varies from the presence of multiple genotypes with no clonal similarity to the presence of only two different clones supra or subgingivally at different sites. While studying natural proportions, it seems that clonal diversity is not informative. To compare similar studies revealed that Streptococcus mutans genotypes showed no pattern of occurrence and number [49].

Consent

All study participants provided informed consent.

Ethical Approval

The study design was approved by the appropriate ethics review board.

Conflicts of Interest

None.

REFERENCES

1. Marsh P, Martin M (2003) Oral Microbiology. Springer 4: 27-55.
2. Armitage GC (1999) Development of a classification system for periodontal diseases and conditions. Ann Periodont 4: 1-6. [Crossref]
3. Burkhardt F, Dahouk S (2009) Microbiological Diagnostics. Bacteriology, Mykology, Virology, Parasitology. Thieme 2: 532-537.
4. Könönen E, Wade WG (2015) Actinomyces and Related Organisms in Human Infections. Clin Microbiol Rev 28: 419-442. [Crossref]
5. Sarkonen N, Könönen E, Summanen P, Könönen M, Joussimies Somer H (2001) Phenotypic identification of Actinomyces and related species.
isolated from human sources. J Clin Microbiol 39: 3955-3961. [Crossref]
6. Lakshmiana Kumar YC, Javherani R, Malini A, Prasad SR (2005) Primary hepatic actinomycosis. Trans R Soc Trop Med Hyg 99: 868-870. [Crossref]
7. Sabbe LJ, Van de Merwe D, Schouls L, Bergmans A, Vaneechoutte M et al. (1999) Clinical spectrum of infections due to the newly described Actinomyces species A. turicensis, A. radingae, A. adhaerens, and A. europaeus. J Clin Microbiol 37: 8-13. [Crossref]
8. Yamane K, Nambu T, Yamakata T, Ishihara K, Tatami T et al. (2013) Pathogenicity of exopolysaccharide-producing Actinomyces oris isolated from an apical abscess lesion. Int Endod J 46: 145-154. [Crossref]
9. Brailsford SR, Tregaskis RB, Leftwich HS, Beighton D (1999) The predominant Actinomyces spp. isolated from infected dentin of active root caries lesions. J Dent Res 78: 1525-1534. [Crossref]
10. Kanellopoulos T, Alexopoulos A, Tanouli ML, Tinakos D, Giannopoulos D et al. (2010) Archimandritis, Primary hepatic actinomycosis. Am J Med Sci 339: 362-365. [Crossref]
11. Dewhirst PE, Chen T, Izard J, Paster BJ, Tanner ACR et al. (2010) The human oral microbiome. J Bacteriol 192: 5002-5017. [Crossref]
12. D’Auto F, Graziani F, Teté S, Gabriele M, Tonetti MS (2005) Periodontitis: from local infection to systemic diseases. Int J Immunopathol Pharmacol 18: 1-11. [Crossref]
13. Costalonga M, Herzberg MC (2014) The oral microbiome and the immunobiology of periodontal disease and caries. Immunol Lett 162: 22-38. [Crossref]
14. Kolenbrander PE (1993) Coaggregation of human oral bacteria: potential role in the accretion of dental plaque. J Appl Bacteriol 74: 795-866. [Crossref]
15. Moutsopoulos NM, Madianos PN (2006) Low-grade inflammation in chronic infectious diseases: paradigm of periodontal infections. Ann N Y Acad Sci 1088: 251-264. [Crossref]
16. Kolenbrander PE, Palmer RJ, Rickard AH, Jakubovics NS, Chalmers Y et al. (1993) Coaggregation of human oral bacteria: potential role in the accretion of dental plaque. J Appl Bacteriol 74: 795-866. [Crossref]
17. Teles FR, Teles RP, Sachdeo A, Uzel NG, Song XQ et al. (2012) Comparison of microbial changes in early redeveloping biofilms on natural teeth and dentures. J Periodontol 83: 1139-1148. [Crossref]
18. Aruni AW, Dou Y, Mishra A, Fletcher HM (2015) The Biofilm Community-Rebels with a Cause. Curr Oral Health Rep 2: 48-56. [Crossref]
19. Halfajee AD, Socransky SS, Patel MR, Song X (2008) Microbial complexes in supragingival plaque. Oral Microbiol Immunol 23: 196-205. [Crossref]
20. Ng HM, Kin LX, Dashper SG, Stuart G, Slakesni K et al. (2016) Bacterial interactions in pathogenic supragingival plaque. Microb Pathog 94: 60-69. [Crossref]
21. Liljemark WF, Bloomquist CG, Bandt CL, Pihlstrom BL, Himrichs JE et al. (1993) Comparison of the distribution of Actinomyces in dental plaque on inserted enamel and natural tooth surfaces in periodontal health and disease. Oral Microbiol Immunol 8: 5-15. [Crossref]
22. Mahajan A, Singh B, Kashyap D, Kumar A, Mahajan P (2013) Interspecies communication and periodontal disease. Sci World J 2013: 765434. [Crossref]
23. Huang R, Li M, Gregory RL (2011) Bacterial interactions in dental biofilm. Virulence 2: 435-444. [Crossref]
24. Vielkind P, Jentsch H, Esrichk R, Rodloff AC, Stingu CS (2015) Prevalence of Actinomyces spp. in patients with chronic periodontitis. Int J Med Microbiol 305: 682-688. [Crossref]
25. Momeni SS, Whudden J, Cheon K, Ghazal T, Moser SA et al. (2016) Genetic Diversity and Evidence for Transmission of Streptococcus mutans by DiversiLab rep-PCR. J Microbiol Methods 128: 108-117. [Crossref]
26. Mougari F, Raskine L, Ferroni A, Marcon E, Sermet Gaudelus I et al. (2014) Clonal relationship and differentiation among Mycobacterium abscessus isolates as determined using the semiautomated repetitive extragenic palindrome sequence PCR-based DiversiLab system. J Clin Microbiol 52: 1969-1977. [Crossref]
27. Babouee B, Frei R, Schultheiss E, Widmer AF, Goldenberger D (2011) Comparison of the DiversiLab repetitive element PCR system with spa typing and pulsed-field gel electrophoresis for clonal characterization of methicillin-resistant Staphylococcus aureus. J Clin Microbiol 49: 1549-1555. [Crossref]
28. Aguadero V, González Velasco C, Vindel A, Gonzalez Velasco M, Moreno J (2015) Evaluation of rep-PCR/DiversiLab versus PFGE and spa typing in genotyping methicillin-resistant Staphylococcus aureus (MRSA). Br J Biomed Sci 72: 120-127. [Crossref]
29. Tenover FC, Gay EA, Frye S, Eells SJ, Healy M et al. (2009) Comparison of typing results obtained for methicillin-resistant Staphylococcus aureus isolates with the DiversiLab system and pulsed-field gel electrophoresis. J Clin Microbiol 47: 2452-2457. [Crossref]
30. Shutt CK, Pounder JI, Page SR, Schaecher BJ, Woods GL (2005) Clinical evaluation of the DiversiLab microbial typing system using repetitive-sequence-based PCR for characterization of Staphylococcus aureus strains. J Clin Microbiol 43: 1187-1192. [Crossref]
31. Venditti C, Vulcano A, D’Azzaro S, Gruber CEM, Sellen M et al. (2019) Epidemiological investigation of an Acinetobacter baumannii outbreak using core genome multilocus sequence typing, J Glob Antimicrob Resist 17: 245-249. [Crossref]
32. Nakamura A, Misawa S, Choman M, Kawakami T, Horii T et al. (2018) Efficacy of PCR-based open reading frame typing assay for outbreak investigation of metallo-β-lactamase-producing Pseudomonas aeruginosa in hematologic unit. J Infect Chemother 24: 1020-1023. [Crossref]
33. Werner G, Fleige C, Neumann B, Bender JK, Layer F et al. (2015) Evaluation of DiversiLab®, MLST and PFGE typing for discriminating clinical Enterococcus faecium isolates. J Microbiol Methods 118: 81-84. [Crossref]
34. Lukinmaa Åberg S, Horsma J, Pasanen T, Mero S, Aulu L et al. (2013) Applicability of DiversiLab repetitive sequence-based PCR method in epidemiological typing of enterohemorrhagic Escherichia coli (EHEC). Food Path Dis 10: 632-638.
35. Stingu CS, Borgmann T, Rodloff AC, Vielkind P, Jentsch H et al. (2015) Rapid identification of oral Actinomyces species cultivated from subgingival biofilm by MALDI-TOF-MS. J Oral Microbiol 7: 2610. [Crossref]
36. Higgins PG, Hujer AM, Hujer KM, Bonomo RA, Seifert H (2012) Interlaboratory reproducibility of DiversiLab rep-PCR typing and clustering of Acinetobacter baumannii isolates. J Med Microbiol 61: 137-141. [Crossref]
37. Avila M, Ojcsus DM, Yilmaz O (2009) The oral microbiota: living with a permanent guest. DNA Cell Biol 28: 405-411. [Crossref]
38. Bergström J (2006) Periodontitis and smoking: an evidence-based appraisal. J Evid Based Dent Pract 6: 33-41. [Crossref]
39. Whittaker CJ, Klier CM, Kolenbrander PE (1996) Mechanisms of adhesion by oral bacteria. *Ann Rev Microbiol* 50: 513-552. [Crossref]
40. Dige I, Raarup MK, Nyengaard JR, Kilian M, Nyvad B (2009) Actinomyces naeslundii in initial dental biofilm formation. *Microbiol* 155: 2116-2126. [Crossref]
41. Liljemark WF, Bloomquist CG, Bandt CL, Pihlstrom BL, Hinrichs JE (1993) Comparison of the distribution of Actinomyces in dental plaque on inserted enamel and natural tooth surfaces in periodontal health and disease. *Oral Microbiol Immunol* 8: 5-15. [Crossref]
42. Zheng H, Xu L, Wang Z, Li L, Zhang J et al. (2015) Subgingival microbiome in patients with healthy and ailing dental implants. *Sci Rep* 5: 10948. [Crossref]
43. Ximénez Fyvie LA, Haffajee AD, Socransky SS (2000) Microbial composition of supra- and subgingival plaque in subjects with adult periodontitis. *J Clin Periodontol* 27: 722-732. [Crossref]
44. Grisold AJ, Zarfel G, Strenger V, Feierl G, Leitner E et al. (2010) Use of automated repetitive-sequence-based PCR for rapid laboratory confirmation of nosocomial outbreaks. *J Infect* 60: 44-51. [Crossref]
45. Koroglu M, Ozpek A, Demiray T, Hafizoglu T, Guclu E et al. (2015) Investigation of clonal relationships of *K. pneumoniae* isolates from neonatal intensive care units by PFGE and rep-PCR. *J Infect Dev Ctries* 9: 829-836. [Crossref]
46. Brolund A, Heggman S, Edquist PJ, Gezelius L, Olsson Liljequist B et al. (2010) The DiversiLab system versus pulsed-field gel electrophoresis: characterisation of extended spectrum β-lactamase producing *Escherichia coli* and *Klebsiella pneumoniae*. *J Microbiol Methods* 83: 224-230. [Crossref]
47. Fontana C, Favaro M, Minelli S, Bossa MC, Testore GP et al. (2008) *Acinetobacter baumannii* in intensive care unit: a novel system to study clonal relationship among the isolates. *BMC Infect Dis* 8: 79. [Crossref]
48. Louws FJ, Rademaker JLM, de Bruijn FJ (1999) THE THREE DS OF PCR-BASED GENOMIC ANALYSIS OF PHYTOBACTERIA: Diversity, Detection, and Disease Diagnosis. *Annu Rev Phytopathol* 37: 81-125. [Crossref]
49. Cheon K, Moser SA, Whiddon J, Osgood RC, Momeni S et al. (2011) Genetic diversity of plaque mutants streptococci with rep-PCR. *J Dent Res* 90: 331-335. [Crossref]