Tobacco-smoking-related prevalence of methanogens in the oral fluid microbiota

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The oral fluid microbiome comprises an important bacterial diversity, yet the presence of archaea has not been reported so far. In order to quest for the presence of methanogenic archaea (methanogens) in oral fluid, we used a polyphasic approach including PCR-sequencing detection, microscopic observation by fluorescence in-situ hybridization, isolation and culture, molecular identification and genotyping of methanogens in 200 oral fluid specimens. In the presence of negative controls, 64/200 (32%) prospectively analysed oral fluid specimens were PCR-positive for methanogens, all identified as Methanobrevibacter oralis by sequencing. Further, fluorescence in-situ hybridization detected methanogens in 19/48 (39.6%) investigated specimens; with morphology suggesting M. oralis in 10 cases and co-infecting Methanobrevibacter smithii in nine cases. M. oralis was cultured from 46/64 (71.8%) PCR-positive specimens and none of PCR-negative specimens; and one M. smithii isolate was co-cultured with M. oralis in one specimen. Multipacer Sequence Typing found one M. oralis genotype per specimen and a total of five different genotypes with 19/46 (41%) of isolates all belonging to spacer-type four. Statistical analyses showed a significant correlation between the PCR-detection of methanogens in oral fluid and tobacco smoking. These data indicate that M. oralis and M. smithii are oral fluid-borne methanogens in tobacco smokers. Both methanogens could be transmitted during intimate contacts such as mother-to-child contacts and kissing.

The repertoire of methanogenic archaea (methanogens) in the oral cavity is limited to six species belonging to the genera Methanobrevibacter (Methanobrevibacter smithii and Methanobrevibacter oralis), Methanosphaera and Methanosarcina. These methanogens have all been detected in subgingival plaque specimens but none of them has ever been documented in the oral fluid. However, it is highly probable that M. oralis could be detected in oral fluid of these patients. Methanobrevibacter oralis is by far the most common methanogen found in this environment with a prevalence of >40%, while other methanogens have been detected in a few studies with a low prevalence of 10–20%. A recent review pooling the data from several studies reported the isolation of M. oralis from the dental plaques of healthy subjects and the identification of M. oralis-like organisms in cases of endodontic infection. Both were detected by polymerase chain reaction (PCR) in periodontitis cases and at peri-implantitis sites. The authors concluded that M. oralis was significantly associated with periodontal disease in terms of abundance when comparing patients and controls, and also diseased and healthy sites within the same patient.

However, methanogens have not been reported in oral fluid of subject free of periodontal disease and based on above mentioned data we formulated the hypothesis that methanogens would be present in oral fluid. In order to test this hypothesis, we undertook a non-randomized, cross-sectional observational study to assess the presence of living methanogens in the oral fluid of voluntary individuals who did not have periodontal disease.

Results and Discussion

In a first step, we investigated methanogens in oral fluid samples using specific PCR-sequencing. Oral fluid was collected at the ostium of the Stenon canal and at the Wharton canal using a previously reported protocol. Of 200 oral fluid specimens here investigated, 64/200 specimens (32%) yielded a positive PCR amplification of
methanogen 16S rRNA gene in the presence of negative controls which all remained negative (Supplementary Table 1). Further sequencing indicated that all the 64 amplicons exhibited a 99% sequence similarity with the reference 16S rRNA gene sequence of \(M. \) oralis strain VD9 (accession NCBI: LN898260.1), \(M. \) oralis strain VD4 (accession NCBI: LN898255.1), \(M. \) oralis strain VD7 (accession NCBI: LN898258.1) and strain \(M. \) oralis VANDE (accession NCBI: LN876656.1).

In a second step, we studied 48 specimens which were in sufficient volume to be investigated by applying fluorescence in situ hybridization (FISH) and direct microscopic examination. While sterile PBS negative controls remained negative, FISH detected methanogens in a total of 19/48 (40%) of investigated specimens including 16 (84.2%) specimens collected from tobacco-smokers (Fig. 1) (Appendix). More precisely, FISH yielded methanogens presenting a bacillary morphology suggesting \(M. \) oralis in 10 cases including eight tobacco smokers and a diplococcus morphology suggesting \(M. \) smithii in nine cases including eight tobacco smokers; for a total of nine cases of co-infection with both methanogens (Fig. 1). These results confirmed that PCR-based data did not merely result from the contamination of the specimens. As the FISH here developed was detecting RNA, these results further suggested that such methanogens were living and not dead ones.

In a third step, we aimed to confirm the viability of methanogens by attempting their isolation and culture. The isolation and culture of methanogens remained negative in negative controls as well as in 136 specimens which were all PCR-negative. However, it was found positive in 46/200 (23%) specimens which were all PCR-positive and 18 PCR-positive specimens remained culture-negative. Identification of colonies yielded \(M. \) oralis in all the culture-positive specimens with 99% sequence similarity with reference 16S rRNA gene sequence of \(M. \) oralis strain VANDE (accession NCBI: LN876656.1) and \(M. \) oralis strain VD9 (accession NCBI: LN898260.1). In addition to \(M. \) oralis colonies, one specimen yielded a second type of colonies identified as \(M. \) smithii with 99% sequence similarity with reference 16S rRNA gene sequence of \(M. \) smithii clone S1-1 (accession NCBI: LK054642.1) (Fig. 2). In order to further characterise these isolates, we genotyped the 46 \(M. \) oralis isolates and our single \(M. \) smithii isolate by using the Multispace Sequence Typing that we previously reported for these two methanogens\(^9,10\). The genotyping of \(M. \) oralis strains revealed the presence of five different genotypes which had been all previously found in dental plaque\(^4\): the genotype 4 was found in 19/46 (41%) specimens while the other four genotypes were found in one to four specimens; and the our single \(M. \) smithii isolate was of genotype 5 previously reported in oral cavity\(^5\).

**Figure 1.** Fluorescence in situ Hybridization (FISH) representative detection of \(M. \) oralis in oral fluid sample. (A) Universal DNA DAPI staining exhibiting blue microorganisms. (B) ARC 915 probe staining the archaeal 16S rRNA gene in red (C) \(mcrA\) probe staining the methanogen \(mcrA\) gene in green (D) overlay of ARC 915 and \(mcrA\) probes exhibiting pink organisms with the diplococcus morphology characteristic of \(M. \) smithii in the yellow square and the morphology of \(M. \) oralis in the green square. Scale bar, 10 \(\mu\)m.
Further anonymous exploitation of the three available clinical variables (age, gender and tobacco smoking) indicated no significant correlation between age ($P$ value $= 0.196$) or gender ($P$ value $= 0.144$) (Fig. 3A) and the prevalence of methanogens in the specimen. However, we observed a highly significant correlation between the presence of methanogens and tobacco smoking ($P$ value $< 0.001$) (Fig. 3B and Table 1).

The prevalence of *M. oralis* as measured by the positivity of PCR, was significantly higher in 47/52 (90%) specimens collected from tobacco smokers than in 17/148 (11%) specimens collected from non-smokers ($P < 0.05$).

In particular, the individual who yielded both *M. oralis* and *M. smithii* in culture was a tobacco-smoker. Further studies will have to determine whether this significant association is just resulting from any here unanticipated confounding factor; or whether tobacco smoking has a direct or indirect biological effect of the viability and growth of methanogens.

We are reporting on the first ever detection of methanogens including *M. oralis* and *M. smithii* in the oral fluid of individuals not suffering from any clinically detectable oral cavity disease. Indeed, the two methanogens here reported are well known inhabitants of the dental plaque and have also been detected in peri-odontitis lesions$^{3,4,11-13}$ as well as in peri-implantitis lesions$^{4,8}$, yet their detection in the oral fluid itself has never been reported. It is now known through the literature that a smoker is likely to have more dental caries due to the increase in dental plaque caused by smoking$^{14}$. We interpreted the data here reported as authentic because the negative controls introduced in every experimental step remained negative and because cross-detection of the methanogens by three different approaches, i.e. FISH, PCR-sequencing and culture was performed.

The PCR-sequencing detection of methanogens was not surprising on the basis of previous observations of methanogens in the oral cavity$^{11,12}$. Indeed, this molecular detection may have merely resulted from bypassing

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**Table 1.** Results of the binomial logistic regression. Null deviance: 250.75 on 199 degrees of freedom. Residual deviance: 134.35 on 196 degrees of freedom. AIC: 142.35NS: No statistically significant; ***$p$-value $< 0.001$.  

|          | Estimate | Standard Error | z value | Statistical significance |
|----------|----------|----------------|---------|-------------------------|
| Intercept| −0.75296 | 1.36493        | −0.552  | NS                      |
| Smoker status | 4.41741 | 0.57504        | 7.682   | ***                     |
| Age      | −0.06295 | 0.04871        | −1.292  | NS                      |
| Sex      | 0.71094  | 0.48670        | 1.461   | NS                      |

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**Figure 2.** *M. oralis* (pale colonies forming a film) and *M. smithii* (white colonies) isolates from an oral fluid specimen collected in a tabacco-smoking individual.

**Figure 3.** Molecular detection of methanogen DNA in oral fluid (A) in males and females and (B) in tobacco smokers and non-smokers.
DNA contamination of the oral fluid from neighbour dental plaque. However, isolation and culture of *M. oralis* and *M. smithii* indicated that in some individuals, some methanogen organisms were living in the oral fluid; and the positivity of FISH reinforced this interpretation by excluding a mere contamination of the specimens by a few methanogens. The data suggest that oral fluid is one component of the oral cavity where methanogens live and that in some individuals, *M. oralis* and *M. smithii* are oral fluid-borne methanogens. Further studies will have to confirm the data here reported; and to determine more precisely the oral fluid component where oxygen-susceptible and hydrogen-requiring methanogens are residing as the oxygen-rich and hydrogen-free extracellular component of oral fluid may not support living methanogens for minutes.

Altogether, the data here reported indicate that methanogens including *M. oralis* and *M. smithii* are common inhabitants of the oral cavity, independently of the presence of any pathology. Accordingly methanogens including *M. oralis* and *M. smithii* are potentially exchangeable between individuals through various occasions of oral fluid direct and indirect exchanges, since oral fluid is one person-to-person exchangeable fluid which may occur during mother-to-child contacts, cross-exchanges of food and fomites, and kissing. During such contacts, living methanogens could be then exchanged and possibly implanted in the receiver's microbiota. This is not without significance since methanogens have been implicated in some diseases of the oral cavity such as periodontitis11-13 and peri-implantitis14 of the digestive tract including obesity15, anorexia16 and chronic constipation17 and were convincingly associated with vaginosis18.

Materials and Methods

Clinical specimens. This cross-sectional study has been approved by the Ethics Committee of the IHU Méditerranée Infection (N°2016-011). In addition, all methods were performed in accordance with the relevant guidelines and regulations. The study was done at the Faculty of Odontology and IHU Méditerranée Infection, both located in Marseille, France. Collection was done between March and June 2017 between 9:00 and 11:00 A.M. in subjects who had not eaten in the past one hour. In order to avoid any bias in the viability of methanogens, oral fluid was collected using a standardized protocol including (1) oral clinical examination by a dentist (E.T., G.A.) comprising of oral mucosae examination, teeth examination including the presence of caries and dental calculus, the presence of oral haemorrhage and oedema; (2) absence of any prior oral disinfection in order to avoid any bias of methanogen collection, (3) collection of freshly emitted oral fluid by two of us with particular expertise buccal cavity anatomy (ET, GA) by gentle rotation of a dry, sterile swab (Deltalab, Carcassonne, France) in the bottom of the mandibular vestibule (part between the dental arch and the gingiva on one side, and the lip and cheek on the other), following a previously validated protocol19 and then under the tongue and the swab was immediately placed into a sterile tube containing a transport medium (TRANS-SAB) as previously described19. Informed consents for study participation have been obtained from all the participants. Were included all adults (>18 years) willing to participate were included. Exclusion criteria included individuals in the incapacity to understand the study, or to sign the informed consent, individuals who declared had implants or periodontal disease. A total of 200 specimens were collected in individuals who had no periodontal disease and no implants after filling in a questionnaire further anonymized, indicating age, gender and the year’s cigarette consumption (Supplementary Text 1). A total of 40 negative control specimens consisted in sterile swabs (used for sampling oral fluid) impregnated with sterile phosphate buffered saline (PBS).

Molecular analyses. After gentle shaking of the TransSAB medium containing the swab, a 250-µL aliquot was added to 0.3-g of acid-washed beads (B106 mm, Sigma, Saint-Quentin Fallavier, France) and to achieve a mechanical lysis, the sample was shaken with a FastPrep BIO 101 apparatus (Qbiogene, Strasbourg, France) at level 6.5 for 2 min. Then, 180 µL of buffer T1 and 25 µL of proteinase K (20 mg/mL) from the NucleoSpin Tissue Mini Kit (Macherey–Nagel, Hoerdt, France) were added. The mixture was incubated overnight at 56 °C. After a second cycle of mechanical lysis, the mixture was incubated for 20 min at 100 °C. Total DNA was then extracted using the NucleoSpin Tissue Mini Kit. Extracted DNA was eluted with 100 µL of elution buffer and the DNA was stored at −20 °C. Extraction of 250 µL of sterile PBS was introduced in each series of DNA extraction as negative control.

A PTC-200 automated thermal cycler (MJ Research, Waltham, USA) was used for the PCRs reaction and a mixture of 50-µL was prepared containing 5 µL of 109 buffer (Qiagen, Courtaboeuf, France) and 0.2 µM of each primer (10 PM) (primers used: 16S rRNA gene was amplified using the broad-range archaeal primers SDArch033aS15 (5′-TCCAGGCCCTACGGG-3′) and SDArch0958aA19 (5′-YCCGGCCGGTTGAMTCACAAAT-3′) (Eurogentec, Seraing, Belgium)); 200 µM (each) of dATP, dCTP, dGTP, and dTTP; 1.5 mM of MgCl2; 1.25 µL of Hot Star Taq polymerase (Qiagen); 13 µL of Dnase/Rnase free distilled water (Gibco, Cergy Pontoise, France) and 5 µL of DNA extracted. The PCR programme was as following: An initial 15-min denaturation step at 95 °C was followed by 40 cycles of denaturation of 30-sec each at 95 °C, a 45-sec annealing step at the appropriate Tm (57 °C or 60 °C) and a 60-sec extension at 57 °C and finally 72 °C for 5 min. For each amplification run, we used distilled water as a negative control. The PCR products were sequenced using the same primers as used for PCRs following this programme: a 1-min denaturation step at 96 °C, followed by 25 cycles of denaturation of 10-sec at 96 °C, a 20-sec annealing at 50 °C and a 4-min extension at 60 °C. The MultiScreen 96-well plates Millipore (Merck, Molsheim, France) containing 5% of Sephadex G-50 (Sigma–Aldrich) was used to purify the sequencing products. The sequences were analysed and edited following the protocol described previously20. We used ChromaPro (http://technelysium.com.au/wp/chromaspro/) and blast programme of NCBI (https://blast.ncbi.nlm.nih.gov/Blast.cgi) to analyse sequences.

Isolation and culture. A total of 250-µL volume of oral fluid sample was seeded in a sterile Hungate tube (Dominique Dutscher, Brumath, France). Hungate tube is a culture tube developed for specifically growing and
maintaining strictly anaerobic bacteria and archaea. It features autoclavable screw cap with a 9-mm opening, a nontoxic, gas impermeable butyl rubber stopper and a disposable screw cap. Each Hungate tube contained 5 mL of SAB broth supplemented with ascorbic acid (1 g/L; VWR International, Leuven, Belgium), uric acid (0.1 g/L; Sigma-Aldrich) and glutathione (0.1 g/L; Sigma-Aldrich). The pH was adjusted to 7.5 with KOH (10 M). 5 mL of SAB broth and oral fluid sample were inoculated with *Bacteroides thetaiotaomicron* (10^10 cells/mL) for hydrogen production as previously described29. The Hungate tube containing the specimen and *B. thetaiotaomicron* in SAB broth was incubated at 37°C with agitation for seven days. Gas chromatography as previously described21 was used to measure the CH4 release by methanogens. We used a Petri dish containing SAB medium supplemented with 15g/L agar and the subculture was seeded in these petri dish and the culture was made according to the protocol previously described20,22. Methanogen colonies appeared after 9-day incubation (average).

**Multiplexer Sequence Typing (MST).** For each specimen five colonies were analysed using protocol as previously described8,10. A PTC-200 automated thermal Cycler (MJ Research) was used for PCRs reaction and followed all the steps described for standard PCR used for molecular analysis of oral fluid specimens. A 25-μL PCR mixture consisting of (2.5 μL of buffer (Qiagen, Courtaboeuf, France), 1 μL of MgCl2, 2.5 μL of 2 mM dNTP, 0.5 μL of each primer 10 μM, 0.25 μL of HotStarTaq DNA Polymerase (Qiagen) and 5 μL of extracted DNA) was prepared and amplified using the following conditions: 95°C for 15 min and 40 cycles at 95°C (30 s), 60°C (30 s) and 72°C (45 s), followed by a 15-min extension at 72°C. For each amplification run, we used distilled water as a negative control. After PCRs reaction, the PCR products were sequenced using the same primers as used for PCRs following this programme: a 1-min denaturation step at 96°C, followed by 25 cycles of denaturation of 10-sec at 96°C, a 20-sec annealing at 50°C and a 4-min extension at 60°C. The MultiScreen 96-well plates Millipore (Merck, Molsheim, France) containing 5% of Sephadex G-50 (Sigma-Aldrich) was used to purify the sequencing products. The sequences were analysed and edited following the protocol described previously9.

**Fluorescence in situ hybridization (FISH).** A subset of randomly selected 48 oral fluid specimens was analyzed by FISH. The oral fluid specimen (400 μL) were taken and washed with PBS two times. The pellet was suspended in 400 μL of PBS and fixed overnight incubation at 4°C with 150 μL of 4% paraformaldehyde. The specimens were centrifuged at 13,000 g for 5 min and the supernatant was removed. Hybridization was performed by using three probes: Archaea-specific Arch915 probe (GTGCTCACCAGCCGCAATTCTCT)24; probe targeting methyl coenzyme M (*mcrA*) which is derived from a PCR primer (TTCAATTGCTAGTTFWGGRTAGTT) prey design by Luton and collaborators24 and negative control was verified by a non-specific probe (ACTTCTACGGGAGGCAGC). Pellet was mixed with 300 μL of 10 μg/mg proteinase K (Qiagen), 20 mM Tris-HCl and 2 mM CaCl2 and incubated for 15 min at 37°C. The pellet was conserved and washed with saline-sodium citrate buffer 2X, then incubated with 150 μL lysozyme (10 mg/mL) (sigma Aldrich) for 15 min at 37°C to be finally washed twice with SSC (2X) and once with sterile distilled water. Detergent treatment using a solution of 0.1% Tween, 0.1% Triton X-100 in PBS was incubated 10 min at room temperature and washing 3 times with a sterile distilled water. Pellet was suspended in 200 μL of sterile distilled water and 8 μL of the solution was fixed in slide by room temperature draying. An ascending series of ethanol (50–80–100%) and air-dried was used to dehydrated the spot. The slide was incubated in addition of 8 μL volume of a hybridization buffer and 5-pmol probe solution at 80°C for 5 min for denaturation and then for hybridization at 46°C for 16 hours. A descending concentration of SSC (X4-X2-X1-X0.5 and X0.2) was used to wash slides and a 5 μL of the universal DNA stain DAPI were added. For the observation, slides were analysed with an epifluorescence microscope Leica DMI6000 with four wavelengths englobing 448FITC nm for reading the mcr1 probe, 545RHOD nm for the 16S DNA stain DAPI were added. For the observation, slides were analysed with an epifluorescence microscope Leica DMI6000 with four wavelengths englobing 448FITC nm for reading the mcr1 probe, 545RHOD nm for the non-specific probe and DAPI.

**Statistical analyses.** All statistical processes were done using the open source statistical language R25. The threshold of 0.05 was the maximal p-value for each statistical conclusion. As our model hypothesis was that presence of *M. oralis* could be associated with the smoker status, taking in account the sex and age influence on consumption, we tested this hypothesis using a binomial logistic regression.

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**Author Contributions**
G.G. performed the PCR and culture experiments and drafted the manuscript. E.T. performed sampling and drafted the manuscript. M.A.B. performed FISH and drafted the manuscript. G.A. performed sampling and drafted the manuscript. H.C. performed statistical analyses and drafted the manuscript. R.R. conceived the study and drafted the manuscript. M.D. conceived the study, interpreted the data and drafted the manuscript. All authors have read and approved the final version of this manuscript.

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