Systemic and Extraradicular Bacterial Translocation in Apical Periodontitis

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Apical periodontitis is an inflammatory disease of microbial etiology. It has been suggested that endodontic bacterial DNA might translocate to distant organs via blood vessels, but no studies have been conducted. We aimed first to explore overall extraradicular infection, as well as specifically by Porphyromonas spp; and their potential to translocate from infected root canals to blood through peripheral blood mononuclear cells. In this cross-sectional study, healthy individuals with and without a diagnosis of apical periodontitis with an associated apical lesion of endodontic origin (both, symptomatic and asymptomatic) were included. Apical lesions (N=64) were collected from volunteers with an indication of tooth extraction. Intracanal samples (N=39) and respective peripheral blood mononuclear cells from apical periodontitis (n=14) individuals with an indication of endodontic treatment, as well as from healthy individuals (n=14) were collected. The detection frequencies and loads (DNA copies/mg or DNA copies/mL) of total bacteria, Porphyromonas endodontalis and Porphyromonas gingivalis were measured by qPCR.

In apical lesions, the detection frequencies (%) and median bacterial loads (DNA copies/mg) respectively were 70.8% and 4521.6 for total bacteria; 21.5% and 1789.7 for Porphyromonas endodontalis; and 18.4% and 1493.9 for Porphyromonas gingivalis. In intracanal exudates, the detection frequencies and median bacterial loads (DNA copies/mL) respectively were 100% and 21089.2 (DNA copies/µL) for total bacteria, 41% and 8263.9 for Porphyromonas endodontalis; and 20.5%, median 12538.9 for Porphyromonas gingivalis. Bacteria were detected in all samples of peripheral blood mononuclear cells including apical periodontitis and healthy groups, though total bacterial loads (median DNA copies/µL) were significantly higher in apical periodontitis (953.6) compared to controls (300.7), p<0.05. Porphyromonas endodontalis was equally detected in both groups (50%), but its bacterial load tended to be higher in apical periodontitis (262.3) than controls (158.8), p>0.05; Porphyromonas gingivalis was not detected. Bacteria and specifically Porphyromonas spp. were frequently detected in endodontic canals and apical lesions. Also, total bacteria and Porphyromonas
INTRODUCTION

Apical periodontitis (AP) is the inflammatory destruction of the apical periodontium as the result of endodontic infection. The hallmark of AP is the development of an osteolytic apical lesion of endodontic origin (ALEO). From an epidemiological point of view, AP represents a highly frequent cause of tooth loss and a non-classic risk factor for several non-communicable diseases (NCD), in part, by inducing low-grade systemic inflammation (Garrido et al., 2019). AP can vary over time between two clinical entities, symptomatic and asymptomatic apical periodontitis (SAP and AAP, respectively), depending on the dynamic balance between bacterial consortia and the host’s response (Hsiao et al., 2012). Among them, the former is considered an immunologically exacerbated stage (Veloso et al., 2020).

ALEOs result from the direct communication between the former sterile pulp tissue and the oral microbiota, often caused by dental caries. The endodontic pathogens organize in multispecies biofilm communities within the root canal, favoring the selection of Gram-negative anaerobic bacteria (Sakko et al., 2016). Though a substantial heterogeneity can be found among geographically diverse populations the black-pigmented anaerobes, *Porphyromonas endodontalis* and *Porphyromonas gingivalis*, are key pathogens in light of their high prevalence within the root canals, prominent virulence factors and/or significance in the bacterial community’s stability and virulence (Róças and Siqueira, 2018).

Bacterial translocation involves the circulation of bacteria and/or their immunogenic products, such as DNA, even without overt infection or clinical signs (Kane et al., 1998; Martinez-Martinez et al., 2009; Ibrahiem et al., 2011). ALEOs might provide a direct pathogenic pathway toward general circulation provoking low-grade systemic inflammation and associated NCD risk. In fact, extraradicular infections have been increasingly identified during the last years, suggesting an initial step in bacterial dissemination beyond the tooth structure, in which they can reach the blood vessels from the apical granuloma (Sunde et al., 2000; Buonavoglia et al., 2013). Furthermore, evidence of endodontic bacterial translocation is indirectly supported by the well-documented enrichment of oral bacterial DNA in diseased peripheral tissues (i.e. atheroma plaques, arthritic joints) (Reichert et al., 2013; Chhibber-Goel et al., 2016; Berthelot and Wendling, 2020). It has been proposed that periodontal bacteria can be transported to distant tissues internalized in immune cells in arthritis (Martinez-Martinez et al., 2009; Ibrahiem et al., 2011), involving mononuclear cell activation, production of pro-inflammatory cytokines, and risk for future NCD complications (Sahingur et al., 2010; Rodriguez-Laiz et al., 2019). Up to now, reports of extraradicular endodontic infection are limited, whereas further bacterial outreach to blood remains unknown. We aimed first to explore overall extraradicular infection, as well as specifically by *Porphyromonas* spp; and their potential to translocate from infected root canals to the blood through peripheral blood mononuclear cells (PBMCs).

MATERIAL AND METHODS

Study Design

Cross-sectional. The study was approved by the Ethics-Scientific Committee of the Central Metropolitan Health Service (N 2017/70) and from the Faculty of Dentistry, Universidad de Chile (N 2016/08). The objectives and procedures of the study were informed to the participants and all of them signed the informed consent or its corresponding forms in the case of underage participants. All procedures followed the ethical standards of the institutional and/or national research committee and with the 1964 Helsinki declaration and its later amendments or comparable ethical standards.

Patient Recruitment

Patients aged between 15 and 40 years old consulting at the dental clinic, School of Dentistry, Universidad de Chile and the Clinic of Surgery, Faculty of Dentistry, Universidad Andrés Bello, Santiago, Chile, were enrolled between 2016 and 2019 if they were otherwise healthy and had a clinical diagnosis of AP in the presence of a radiographic apical radiolucency due to extensive caries in a tooth with negative sensitivity pulpal test with no previous endodontic intervention. SAP and AAP were diagnosed when clinical symptoms in response to percussion were present or absent, respectively, according to previously defined diagnostic terminology (Gutmann et al., 2009). Controls met the same criteria and the absence of any tooth with ALEOs. Exclusion criteria were obesity (body mass index ≥30 kg/m²), pregnancy, moderate to severe periodontitis, and antibiotic and/or anti-inflammatory drug consumption 3 months before the study (Garrido et al., 2019).

Sample Collection

Periapical tissue samples were collected from individuals having indication of tooth extraction. Tissue samples from ALEOs (n=64), including AAP (n=29) and SAP (n=35), or healthy periodontal ligaments (HPL) as negative controls (n=9), were obtained by surgical separation of the tissue from the root surface with sterile curettes. Then the samples were washed with 3 mL of sterile NaCl solution before their storage in 100 μL of RNAlater (Qiagen, Valencia, CA, USA) at -80°C until processed for tissue homogenization.
Root canal exudates and the respective blood samples were collected from AP patients having indication of endodontic treatment. For intracanal exudate sampling (N=39; AAP=29 and SAP=10), the involved tooth was isolated with a rubber dam and disinfected with 70% ethanol solution. After coronal access, sterile paper points (Maillefer®) were inserted into the infected root canal for 1 minute, transported in a vial containing 1 mL of cold sterilized reduced transport fluid (RTF), and stored at -80°C for posterior analysis. Blood samples were collected from a subset of AP patients (n=14) and also from healthy control volunteers (n=14) by venipuncture of the antecubital vein (Fouad et al., 2002; Rocas and Siqueira, 2011). PBMCs were isolated using Ficoll-Paque Premium 1.073 (GE Healthcare®), following the manufacturer’s instructions.

DNA Extraction
Apical tissue samples were homogenized in a lysis buffer with lysozyme 20 mg/mL at 37°C for 1 h. DNA was extracted using a commercial kit to obtain bacterial DNA (NucleoSpin® TriPrep, Macherey-Nagel), according to the recommendations of the manufacturer. Intracanal exudate DNA was extracted by the boiling modified method (Queipo-Ortuño et al., 2008); the Eppendorf tubes containing the paper points and the RTF medium were boiled in a water bath for 10 minutes, followed by cooling down to -20°C for 10 minutes and were finally centrifuged for five minutes at 1000 rpm. PBMC bacterial DNA was extracted in a lysis buffer using lysozyme followed by the TE buffer and DNEasy isolation kit (QIAGEN Inc., Valencia, CA, USA), according to the manufacturer. The concentration and quality of DNA were confirmed by the 260:280 ratio in a spectrophotometer (Bio-Tek, Winooski, VT).

qPCR Assay
Quantitative PCR (SteponePlus®; Applied Biosystems, Singapore) was carried out to determine bacterial loads and frequencies of detection of total bacteria, P. gingivalis and P. endodontalis. Primer sets are shown in Table 1. Briefly, total bacteria were quantified using specific primers targeting 16S ribosomal RNA (rRNA) gene previously published (Nadkarni et al., 2002); as well as P. gingivalis, which was identified by using previously validated 16S rRNA gene primers (Byrne et al., 2009). The primer set for targeting P. gingivalis included 16S rRNA from base 729 to 1192 (GenBank: L16492.1). For P. endodontalis identification, specific primers targeting the heat shock protein 60 were used (Hsp60 gene) (Sakamoto and Ohkuma, 2010). The Hsp60 gene sequence was obtained from GeneBank NIH genetic sequence database (GenBank: AB547580.1). Primer pairs were designed and assessed manually in the Primer Blast tool, considering the real-time PCR guide (BIO RAD). Which included having a melting temperature (Tm) between 50°C and 65°C, a GC content of 50–60%, to be limited to obtain an amplicon ranging from 75 to 200 bp. The selected primer pairs, targeting from base 84 to 201, were checked for specificity and cross-reactivity using Primer Blast and UniProt databases. Experimental validation included: firstly, the analysis of the molecular weight of the DNA fragments of three positive samples for P. endodontalis in agarose gel electrophoresis. Secondly, positive samples in step 1 were confirmed by Sanger sequencing (Macrogen, Seoul, Korea), showing 100% alignment with P. endodontalis (Hsp60 gene). Also, to determine the primer specificity, a dissociation curve analysis was performed, using temperatures between 60°C and 95°C for the recognition of non-specific product formation or contamination.

Each qPCR was performed using KAPA SYBR® Fast qPCR Kits (KAPA Biosystems, Woburn, MA, USA). Positive controls consisting of P. gingivalis ATCC 33277 and P. endodontalis ATCC 35406 DNA, and negative control (no DNA) were included in each experiment. Cycling conditions consisted of the following steps: 95°C for 3 min, 40 cycles: total bacteria 95°C for 15 sec and 60°C for 1 min; P. gingivalis 95°C for 3 sec and 58°C for 1 min; P. endodontalis 95°C for 30 seconds and 58°C for 1 min. As it was stated above, both in silico and experimental approaches were used for primers validation.

Absolute quantifications of total bacterial and individual species loads were carried out by comparing the threshold cycle (Dorn et al., 2002) values of the test samples to standard curves of known DNA copy numbers of P. gingivalis and P. endodontalis (Supplementary Figure 1). The lower and higher detection limits were 10^2 and the 10^8 DNA copy numbers, respectively. All experiments were performed with a linear quantitative detection range, Supplementary Figure 1. Bacterial loads were expressed as DNA copies/µL in the case of ALEOs, and DNA copies/µL in the case of intracanal exudates and PBMCs.

Statistical Analysis
The sample size was calculated during a pilot study based on the positive detection of P. endodontalis in AAP versus SAP in ALEOs. Considering an 80% statistical power and 5% alpha error, a minimum of 23 samples in each group (SAP and AAP) were required.

Data distribution was evaluated by the Shapiro-Wilk test. Fisher’s exact Chi-square tests were used to compare the frequency of bacterial detection between two groups; Mann-Whitney test was used to compare bacterial loads between two groups; and McNemar’s and Spearman’s correlation tests were conducted to compare the associations in the frequencies of detection and bacterial loads, respectively between intracanal

| Target bacteria | Target gene | Forward primer (5'-3') | Reverse primer (5'-3') |
|-----------------|-------------|------------------------|-----------------------|
| Total Bacteria  | 16S rRNA    | TCCTACGCGAGGACGAGGT    | GGACTACCAAGGGTATCTAATCTGT |
| P. endodontalis | Hsp 60      | TATTGCAAGGCTTGGCTAACC  | TTCTCTGCTGCCCCATTAGCAGG  |
| P. gingivalis   | 16S rRNA    | AGGCAGCCTGGCATACTGCG   | ACTGTTAAGTAACTACGATGT    |

**TABLE 1** | Primers used to determine the bacterial loads and frequency of target bacteria.
exudates and PBMCs within AP individuals. Analyses were performed using STATA 12® (StataCorp LP, TX, USA). The level of statistical significance was p<0.05.

RESULTS

This study included a total of 126 individuals. Demographics and smoking habits are shown in Table 2. Figures 1A, B shows extraradicular infection rates for total bacteria, P. endodontalis, and P. gingivalis. The detection frequency, expressed as n (%), and bacterial loads expressed as medians (interquartile range) in ALEOs were 46 (70.8%) and 4521.6 (7789.1) DNA copy number/mg, respectively for total bacteria; 14 (21.5%) and 1789.7 (1338.8) DNA copy number/mg for P. endodontalis; 12 (18.4%) and 1493.9 (26919.9) DNA copy number/mg for P. gingivalis. A higher frequency of detection was identified for P. endodontalis in symptomatic ALEOs compared to asymptomatic ALEOs (p = 0.003), while no significant differences were found in the detection frequencies of total bacteria and P. gingivalis or bacterial loads (p > 0.05). Healthy periodontal ligaments confirmed no bacterial detection.

The frequency of detection and loads of total bacteria and Porphyromonas spp. in intracanal exudates are shown in Figures 2A, B. The frequencies of detection expressed as n (%) and bacterial loads expressed as median (interquartile range) DNA copy number/µL, respectively in AP were n=39 (100%) and 21089.2 (263450.7) for total bacteria; n=16 (41%) and 8263.9 (85001.8) for P. endodontalis; and n=8 (20.5%), 12538.9 (72957.3) for P. gingivalis. The frequency of detection and bacterial loads for P. endodontalis were 48.3% and 6127.9 (121214.7) in AAP, and 20%, and 9331.2 (6437.1) in SAP exudates, (p > 0.05). P. gingivalis was only identified in intracanal exudates from AAP with positive detection of n=8 (27.6%) and a bacterial load of 12538.9 (107919.3) DNA copy number/µL.

The frequency of bacterial DNA detection and loads in PBMCs are presented in Figures 3A, B. Bacteria were detected in 100% of the PBMC samples from AP and control groups. The total bacterial load was significantly higher in AP individuals compared to healthy controls (953.6 vs 300.6 median DNA copy number/µL respectively; p = 0.0002). On the other hand, P. endodontalis was detected in 50% of the samples of both, AP individuals and controls, showing an even higher frequency of detection than in intracanal exudates and apical lesions, though its bacterial loads were low, and no statistically significant differences were found between AP and healthy groups (p > 0.05). P. gingivalis was not detected in any of the evaluated PBMC samples (data not shown).

To explore the potential endodontic origin of the bacteria detected in PBMCs, the frequencies and loads of total bacteria, P. endodontalis, and P. gingivalis were associated between the intracanal samples and PBMCs within each AP individual (Table 3). Detection of total bacteria was positive in all intracanal exudates and PBMC samples (14/14). P. endodontalis was identified in only two intracanal exudate samples (2/14) and seven PBMC samples (7/14) in AP individuals. From them, only one patient was positive for P.

| TABLE 2 | Demographic data and smoking habits of study groups. |
| Parameters | Tissue Samples | Intracanal (n = 39) | PBMCs |
| | ALEOs (n = 64) | HPL (n = 9) | AP (n = 14) | Control (n = 14) |
| Age [years, median (IQR)] | 38 (22.5) | 22 (5) | 25 (5) | 24 (9.25) | 22 (6.5) |
| Females (n, %) | 27 (42.2%) | 6 (66.7%) | 18 (46.2%) | 5 (28.6%) | 4 (28.6%) |
| Smokers (n, %) | 28 (44.4%) | 2 (22%) | 14 (35%) | 9 (64.3%) | 1 (7.1%) |
| Educational level (median) | Full high school | Full high school | Full high school | Full high school | Full high school |

ALEOs, Apical lesions of endodontic origin; HPL, healthy periodontal ligament; PBMCs, Peripheral blood mononuclear cells; IQR, interquartile range.

**FIGURE 1** | (A) Frequency of detection and (B) copy numbers of 16S rRNA gene of total bacteria and P. gingivalis, and Hsp60 gene of P. endodontalis in total ALEOs (n = 64) including AALEOs (n = 29) and SALEOs (n = 35). ALEOs Apical lesions of endodontic origin; AALOE asymptomatic apical lesions of endodontic origin; SALEOs symptomatic apical lesions of endodontic origin. *p < 0.05.
**endodontalis** at both, the intracanal exudate and the respective PBMCs. On the other hand, *P. gingivalis* was detected in three intracanal samples, though it was not detected in PBMCs. Regarding bacterial loads, no correlation was found between intracanal exudates and PBMCs (Spearman’s rho = 0.28, p = 0.33; data now shown).

**DISCUSSION**

ALEO is the result of a polymicrobial infection of the root canal system that causes local inflammation and destruction of the periapical tissues. Though bacteria have been largely assumed to be confined within the necrotic tooth canal, emerging evidence suggests that they can disseminate beyond the tooth structures (Liljestrand et al., 2016). Based on bacterial DNA detection, in the current study, we show a high frequency of extraradicular infection. Specifically, *P. endodontalis* was detected with a higher frequency in symptomatic compared to asymptomatic ALEO. Moreover, endodontic bacteria were detected in circulating mononuclear cells, whereas total bacterial loads were higher in PBMCs from AP compared to healthy individuals.

The presence of bacteria in the root canal system of teeth with AP has long been documented (Ricucci and Siqueira, 2010), but the concept of extraradicular infection is rather recent with only a few studies available. Given their high sensitivity, molecular techniques are widely being used to characterize the microbial communities (Rôças and Siqueira, 2008; Zhang et al., 2010; Rôças et al., 2011). Herein, we seek for unspecific oral bacteria and specific *Porphyromonas* spp. in ALEOs by qPCR. Identification of bacterial DNA in PBMCs and intracanal exudates was performed by qPCR.
of viable Porphyromonas spp. was formerly demonstrated through bacterial culture techniques in periapical abscesses (Van Winkelhoff et al., 1985). As part of the endodontic microbiome, Gram-negative species, such as P. endodontalis and P. gingivalis, are likely to have a clinical significance due to their wide repertoire of virulence factors (Sundqvist et al., 1989).

Our results revealed high detection frequencies of total bacteria (70.8%), P. gingivalis (18.4%), and P. endodontalis (21.5%) in ALEOs. Specific identification of P. endodontalis and P. gingivalis with frequencies of 45% and 27% respectively, was also reported in ALEOs from persistent apical periodontitis based on qPCR (Zhang et al., 2010). The current study also explored the association between extraradicular infection and clinical symptoms, reporting a significantly higher detection frequency of P. endodontalis in ALEOs from SAP compared to AAP, while the detection frequencies and bacterial loads of P. gingivalis and total bacteria did not show differences. Though experimental mono-infection by P. endodontalis generates low pathogenicity compared to P. gingivalis, anaerobic mixed communities containing both species can trigger severe infective responses (Van Winkelhoff et al., 1985). Presumably, P. endodontalis might contribute to the bacterial mixed community to become more virulent, recruiting an enhanced immune response and giving rise to clinical symptoms. Therefore, it seems plausible that extraradicular translocation of P. endodontalis plays a relevant role in active forms of ALEOs.

The infection of the root canal system has been extensively characterized in AP (Garbia et al., 1994; Fouda et al., 2002; Siqueira, 2002; Ricucci and Siqueira, 2010). Bacteria have been detected up to 100% within root canals from primary endodontic infection by molecular techniques (Conrads et al., 1997; Rôças and Siqueira, 2010). In line with this, intracanal samples of AP were all positive for bacterial DNA in our study. In earlier studies, the intracanal detection of Porphyromonas spp. revealed low recovery rates (9-23.9%) by culture methods (Hashioka et al., 1992; Baumgartner et al., 1999; Jacinto et al., 2003), though recent data based on molecular techniques report higher prevalence in primary endodontic infections (between 23.3-50% and 33-44%, respectively) (Tomazinho and Avila-Campos, 2007; Cao et al., 2012). Similarly, our results show detection frequencies of 41.0% for P. endodontalis and 20.5% for P. gingivalis in intracanal samples, confirming the relevant etiologic role of these black-pigmented anaerobic rods in AP (Dougherty et al., 1998; Baumgartner et al., 1999; de Oliveira et al., 2000).

Though the etiologic role of the intracanal bacterial infection is a matter of fact, its association with clinical symptoms remains unclear. Our results show that the detection frequencies and loads of total bacteria and Porphyromonas spp. in endodontic canals remained similar among AAP and SAP. Previous studies have reported comparable detection rates in root canals, especially for P. endodontalis, varying between 56% to 62% in AAP; and 40% to 69% in SAP (Van Winkelhoff et al., 1985; Rôças et al., 2002; Rôças and Siqueira, 2008; Rôças et al., 2011; Cardoso et al., 2016); while the available evidence of the detection of P. gingivalis ranges between 8-38% in SAP and 4-29% in AAP (Wang et al., 2010; Buonavoglia et al., 2013). The frequent intracanal detection of Porphyromonas spp. in association with AAP might be related to the confinement of the infection to the root canal. Importantly, in this condition bacteria gain no access to blood vessels as pulp tissue is necrotic (Rôças and Siqueira, 2010). Moreover, our results showed that the relative detection of both, P. endodontalis and P. gingivalis, was higher in ALEOs than intracanal exudates from SAP; and conversely, they were higher in intracanal exudates than ALEOs in AAP, even though a direct comparison cannot be performed because ALEOs and intracanal exudates were obtained from different individuals based on their treatment indications, altogether these antecedents suggest that the transit to an acute periapical process and the consecutive onset of clinical symptoms might rely on the host’s inability to confine the endodontic pathogens -such as P. endodontalis- within the root canals and extraradicular infection might represent the first step for bacterial dissemination.

Given the anatomic relation of ALEOs with the bloodstream, it is plausible that inflammatory mediators, bacteria, and/or their products translocate from their endodontic sites to the systemic circulation (Garrido et al., 2019). Oral bacteria can be transported to distant tissues as free circulating DNA and/or internalized in immune cells (Martinez-Martinez et al., 2009; Ibrahim et al., 2011), though the former is less probable because circulating oral bacteria are cleared within few minutes (Minasyan, 2019). Accordingly, we screened for bacterial DNA within PBMCs from AP and healthy individuals and found significantly higher total bacterial loads in the former, suggesting that ALEOs create a favorable environment for the overgrowth of microorganisms and their systemic dissemination. Moreover, P. endodontalis DNA was also detected in PBMCs from individuals in both groups (50%). In fact, a previous study reported the carriage of bacterial DNA (16S rDNA) in PBMCs from arthritis and control individuals (Ibrahim et al., 2011). Despite no studies are available in AP, a pathophysiological role for blood dendritic cells in systemic dissemination of periodontal pathogens to atherosclerotic plaques was proposed, particularly of P. gingivalis (Carrion et al., 2012). Herein we failed to detect P. gingivalis in PBMCs, probably because our study individuals were free from periodontitis. Instead, P. endodontalis might analogously translocate to distant sites from oral/endodontic sites via PBMCs and has also the ability to invade endothelial cells in vitro (Dorn et al., 2002). Additionally, bacterial carriage has been associated with mononuclear cell activation, production of pro-inflammatory cytokines, prolonged survival of circulating monocytes, and further risk of future complications of systemic non-communicable diseases, including cirrhosis and hemodialysis (Hazzah et al., 2015; Rodriguez-Laiz et al., 2019). These results are in line with the higher inflammatory burden of serum markers reported in patients with ALEOs (Garrido et al., 2019). Altogether, these preliminary studies support that oral/endodontic microorganisms can translocate from their habitats to systemic circulation and reach distant organs using mononuclear cells as Trojan horses. The capacity for intracellular survival of bacteria within phagocytes is likely a critical factor facilitating the dissemination. Some bacteria can hide within autophagosomes, where they protected themselves from phagocytic killing (Berthelot and Wendling, 2020).
Despite the abovementioned bacterial translocation in AP, no association was found between the presence and loads of bacteria between endodontic canals and PBMCs within the study individuals. Oral bacteria and specifically *P. endodontalis* might coexist in various oral habitats, including subgingival sites and periodontal pockets, as well as oral mucosa (i.e. tonsils and tongue), where they can also be phagocytosed by mononuclear cells and reach the general circulation (Lombardo Bedran et al., 2012; Scapoli et al., 2015).

Although surgical cultures are considered the reference method to determine the presence of microorganisms (Zweitzig et al., 2013), molecular analysis is more sensitive, especially in anaerobic conditions, such as necrotized root canals (Baumgartner, 2004). Particularly, *Porphyromonas* is considered fastidious genera and culture methods may underestimate their frequency (Gomes and Herrera, 2018). Also, no viable oral bacteria can be recovered from peripheral lesions in most studies (Martinez-Martinez et al., 2009; Chhibber-Goel et al., 2016), whereas PCR has proven to be highly sensitive to detect bacterial translocation (Kane et al., 1998). Instead, bacterial DNA is much more likely to disseminate from oral niches through blood and can trigger and perpetuate inflammation after bacterial clearance.

Summarizing, total bacteria and specific *Porphyromonas* spp. were frequently found in endodontic canals and extra-radicular infection in AP patients. Extraradicular bacteria and specifically, detection of *P. endodontalis*, associated with asymptomatic forms of the disease and might represent a first step in the dissemination process. Also, *P. endodontalis* was frequently detected in PBMCs from study individuals and higher total bacterial loads were found in AP, supporting a role for these cells in the bacterial carriage during AP.

**DATA AVAILABILITY STATEMENT**

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

**ETHICS STATEMENT**

The studies involving human participants were reviewed and approved by Ethics-Scientific Committee of the Central Metropolitan Health Service (N 2017/70) and from the Faculty of Dentistry, Universidad de Chile (N 2016/08). Written informed consent to participate in this study was provided by the participants’ legal guardian/next of kin.

**AUTHOR CONTRIBUTIONS**

MH, MG, AH: Conceptualization and design. MG, MB: Data curation. MH: Funding acquisition. MH, AF, MB, AH: Formal analysis. MB, JA, AH: Laboratory experiments. MH: Project administration. AH: Design the figures. MH: Resources. MH: Software. MG, MH: Supervision. MB, AF, MH: Writing manuscript. All authors contributed to the article and approved the submitted version.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fcimb.2021.649925/full#supplementary-material

**Supplementary Figure 1 | Quantitative PCR standard curves obtained for total bacteria (A), *P. gingivalis* (B) and *P. endodontalis* (C) quantification. The regression line from the 10-fold dilutions curve was used to determine the copy number of all samples. All reactions were performed in duplicate.**

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Conflict of Interest: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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