Polymicrobial periodontal disease triggers a wide radius of effect and unique virome

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Periodontal disease is a microbially-mediated inflammatory disease of tooth-supporting tissues that leads to bone and tissue loss around teeth. Although bacterially-mediated mechanisms of alveolar bone destruction have been widely studied, the effects of a polymicrobial infection on the periodontal ligament and microbiome/virome have not been well explored. Therefore, the current investigation introduced a new mouse model of periodontal disease to examine the effects of a polymicrobial infection on periodontal ligament (PDL) properties, changes in bone loss, the host immune response, and the microbiome/virome using shotgun sequencing. Periodontal pathogens, namely Porphyromonas gingivalis, Treponema denticola, Tannerella forsythia, and Fusobacterium nucleatum were used as the polymicrobial oral inoculum in BALB/cByJ mice. The polymicrobial infection triggered significant alveolar bone loss, a heightened antibody response, an elevated cytokine immune response, a significant shift in viral diversity and virome composition, and a widening of the PDL space; the latter two findings have not been previously reported in periodontal disease models. Changes in the PDL space were present at sites far away from the site of insult, indicating that the polymicrobial radius of effect extends beyond the bone loss areas and site of initial infection and wider than previously appreciated. Associations were found between bone loss, specific viral and bacterial species, immune genes, and PDL space changes. These findings may have significant implications for the pathogenesis of periodontal disease and biomechanical properties of the periodontium. This new polymicrobial mouse model of periodontal disease in a common mouse strain is useful for evaluating the features of periodontal disease.

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INTRODUCTION

Periodontitis is an inflammatory disease triggered by a microbial dysbiosis that affects the supporting tissues around teeth and it eventually leads to tooth loss if left untreated. Periodontitis has been associated with an increased risk of diabetes mellitus, cardiovascular diseases, adverse pregnancy outcomes, respiratory infections, and rheumatoid arthritis1,2. A dysbiotic oral microbiota is the initiating factor in the pathogenesis of periodontitis that leads to a dysregulated host immune response3,4. There are more than 700 species colonizing the oral cavity5 and hundreds of those can be present within oral biofilms6,7. Among the biofilm-associated microbiota, several bacterial species are highly associated with periodontal disease; namely Porphyromonas gingivalis, Treponema denticola, and Tannerella forsythia. This pathogenic consortium of microbes is known as the “Red Complex” due to their strong association with clinical parameters of severe periodontal disease, such as periodontal pocket depths and bleeding on probing8–11. However, recent studies using high throughput sequencing approaches are shedding new light on novel bacterial signatures and a larger and more diverse microbiome associated with periodontal disease12,13. Another gram-negative bacterium, namely Fusobacterium nucleatum, which belongs to the “orange complex” and is closely related to the red complex, is also associated with periodontitis8. F. nucleatum can coaggregate with a significant number of bacteria in the oral cavity and is thought to be an important “microbial bridge” during dental plaque formation14,15. These pathogenic bacteria contribute to periodontal disease via a variety of mechanisms, including through secretion of proteolytic enzymes or proteases16, activation and modulation of the host immune response via bacterial LPS and other surface effector molecules17,18, and invasion of host cells19. These molecular and cellular processes lead to destruction of the periodontal supporting tissues.

Although viruses are also present within the oral cavity with differences observed between health and disease states, there is limited knowledge about their relationship to periodontal disease20. Studies suggest that viruses, including herpes simplex virus-1,2, Epstein–Barr virus, and cytomegalovirus play important roles in periodontal disease21–29. Also, in recent studies, other different viruses have been associated with periodontal disease, however, a comprehensive evaluation of the oral virome in relationship to periodontal disease using shotgun sequencing has not been carried out in mice and to a limited extent in humans30–33.

The effects of a defined polymicrobial infection on periodontal ligament properties and on the microbiome/virome have not been examined. Thus, in this study, a new polymicrobial mouse model of periodontal disease, induced by oral infection with Porphyromonas gingivalis, Treponema denticola, and Tannerella forsythia, was used to examine the effects of a polymicrobial infection on periodontal ligament (PDL) properties, changes in bone loss, the host immune response, and the microbiome/virome using shotgun sequencing.
P. gingivalis, T. denticola, T. forsythia, and F. nucleatum in a common mouse strain, was introduced to examine the effects on periodontal ligament properties and the microbiome/virome.

RESULTS
A polymicrobial oral infection can be successfully obtained in a common mouse strain
A polymicrobial infection model of periodontal disease in a commonly employed mouse strain (BALB/cByJ) was used to evaluate periodontal bone loss and related parameters (Fig. 1). A polymerase chain reaction (PCR)-based approach was used to confirm oral infection with all four periodontal pathogens (P. gingivalis, T. denticola, T. forsythia, and F. nucleatum) in this strain of mice (Fig. 2; Table 1). The oral swab results indicated that all four bacteria were detectable throughout the experimental period from 1 to 8 weeks post infection; except T. denticola, which was always present except at 4 weeks. All pathogens were present at higher levels in the infected versus control group for all time points. These differences were statistically significant for all pathogens except T. denticola, which showed a trend toward higher levels compared to controls. The highest levels of the four pathogens were detected at 1 week post infection, except for T. forsythia, which showed even higher levels at 8 weeks. T. denticola was not detected at 4 weeks in either infection or control group and at 8 weeks it was only detected in the infection group. The number of mice exhibiting infection with the different pathogens varied. For example, P. gingivalis was present in all mice at all time points. However, T. denticola was present in generally few mice throughout the study period. T. forsythia and F. nucleatum were similar to P. gingivalis in that they were present in most mice throughout the study.

The polymicrobial infection induces a host antibody response against all periodontal pathogens
To further validate the infection and to evaluate the systemic host response to the polymicrobial infection, serum antibody levels to the four periodontal pathogens were evaluated using an ELISA approach (Fig. 3). Infected mice showed a significant IgG antibody response to all four periodontal pathogens compared to the uninfected control mice. Specifically, the IgG antibody levels for P. gingivalis were 10,279.13 ng/ml ± 845.68 in the infected group compared to 257.37 ng/ml ± 87.18 in the control. The IgG antibody levels for T. denticola were 10,395.19 ng/ml ± 975.65 in the infected group compared to 218.21 ng/ml ± 25.76 in the control. The IgG antibody levels for F. nucleatum were 17,702.02 ng/ml ± 614.57 in the infected group compared to 310.64 ng/ml ± 27.19 in the control. The IgG antibody levels for T. forsythia were 16,848.83 ng/ml ± 1509.53 in the infected group compared to 367.21 ng/ml ± 41.82 in the control.

The polymicrobial infection induces a significant change in immune and cytokine gene expression
To further characterize this polymicrobial mouse model, immune gene profiling was performed on the gingival tissues of the infected and control mice. Using an RNA array of 84 genes, this profiling revealed 18 genes that were differentially altered 1.5-fold or greater in the infected mice compared to the control uninfected mice (Table 2), although p values were greater than 0.05. The most differentially elevated genes, included secretoglobin gene super family cytokine (8.473-fold change) and interferon gamma-inducing factor (2.287-fold change), and the most differentially decreased gene was tumor necrosis factor ligand 1C (~ 2.336-fold change). The Scgb3a1 gene encodes a small secreted protein that may function in the inhibition of cell growth. Expression of the related gene in humans is downregulated via DNA methylation in breast cancers and other tumor types. Alternatively, spliced transcript variants encoding multiple isoforms have been observed for this gene.

Alveolar bone loss and intrabony defects are significantly increased with a polymicrobial infection
Once the local infection with and systemic response to the four periodontal pathogens was confirmed, the alveolar bone response to the polymicrobial infection was evaluated in the mice. Bone loss was evaluated in the maxillary and mandibular molars of the mice. After 8 weeks of inoculation and infection with the periodontal pathogens, mice exhibited significantly more alveolar bone loss (0.136 mm ± 0.011) compared to the control group (0.088 mm ± 0.004) (Fig. 4a, b).

The presence of alveolar intrabony defects was also evaluated following infection. Infected mice showed significantly more intrabony defects (42/72 or 58% of sites) compared to control uninfected mice (14/72 or 19% of sites) (Fig. 4c).

The polymicrobial infection induces a widening of the periodontal ligament space throughout, thereby inducing a broad radius of effect
Given that the role of periodontal pathogens in modulating periodontal ligament parameters has not been previously examined, this study evaluated the dimensions of the periodontal ligament space following infection with the four periodontal pathogens. The polymicrobial infection triggered a widening of the periodontal ligament space throughout its length as measured by micro-CT analysis (Fig. 5). Specifically, the infected group exhibited significantly increased periodontal ligament space/width (45 μm ± 1) compared to the control group (37 μm ± 2). Notably, the widened periodontal ligament space was detected throughout the periodontal ligament and even at a great distance from the site of infection (gingival margin) and bone loss thereby inducing a broad radius of effect.

The polymicrobial infection induces significant changes in viral diversity and the virome
For the first time, a metagenomic shotgun sequencing approach was used to examine the microbiome and virome of periodontal disease in a mouse model with a defined polymicrobial infection. Analysis of the gingival microbiome/virome data revealed that there is a significant increase in viral content and diversity in

Fig. 1 Mouse infection and oral sample collection timeline. Eight polymicrobial infection sessions were carried out once per week from the 3rd to the 10th week. Oral swab samples were collected on the day before antibiotic treatment, the day before infection and at 1, 4, and 8 weeks following the initial infection. Blood and tissue specimen collection was performed at euthanasia following 8 weeks of infection.
response to the polymicrobial infection in mice (Fig. 6). Although, the bacterial content and diversity was not significantly different. Specifically, the polymicrobial infection group had similar bacterial reads (panel a, p value 0.92) but significantly more virus reads (panel b, p value = 0.04) compared to the control group. In addition, the infection group had a slight decrease in bacterial diversity (panel c, p value = 0.72) and a slight increase in viral diversity (panel d, p value = 0.18) compared to the control group.

Specific viral and bacterial species are associated with bone loss and changes in PDL space

Specific viral and bacterial species were associated with bone loss in mice with the polymicrobial infection (Fig. 7a–h; Table 3). Specifically, genus Marinobacter (false discovery rate (FDR) = 0.097, $R^2 = 0.62$, decreases with bone loss), genus Gammaretrovirus (FDR = 0.065, $R^2 = 0.71$, increases with bone loss), genus Pasteurella (FDR = 0.249, $R^2 = 0.45$, decreases with bone loss), genus Camphylobacter (FDR = 0.213, $R^2 = 0.50$, decreases with bone loss), species Porcine type-C oncovirus (FDR = 0.109, $R^2 = 0.60$, increases with bone loss), species Bat gammaretrovirus (FDR = 0.069, $R^2 = 0.69$, increases with bone loss), species Marinobacter sp B9-2 (FDR = 0.109, $R^2 = 0.62$, decreases with bone loss), and Golden hamster intracisternal A particle H18 (FDR = 0.069, $R^2 = 0.73$, increases with bone loss) were associated with bone loss.

The species Porcine type-C oncovirus (FDR = 0.032, $R^2 = 0.96$, increases with peak width) was associated with increased PDL space or (peak width) (Fig. 7i).

In addition, we studied the relationship between the microbial environment and the immune response and found some weaker associations between bacteria/virus species and immune genes (Supplemental Fig. 1). The species Gibbon ape leukemia virus (FDR = 0.2602, $R^2 = 0.78$, increases with gene-expression level) was associated with the immune gene Tnfsf14 and the species Clostridiales bacterium 1 7 47FAA) was associated with the immune gene Adipoq (FDR = 0.2602, $R^2 = 0.88$, increases with gene-expression level). Also, there were weaker associations between bacteria/virus species and PDL space (Supplemental Fig. 2).

**DISCUSSION**

Several different mouse models of periodontal disease that mimic the human condition have been proposed in the literature, including ligature models, injection models, mono-infection models inoculated with only one strain of bacteria, and even polymicrobial infection models. Ligature and injection models develop aggressive lesions, and the destruction of periodontal tissues is always localized and rapid. These characteristics do not mimic the chronic nature of periodontal disease. Furthermore, mono-infection models do not recapitulate the polymicrobial nature of periodontal disease. Several studies have suggested that the host immune and physiologic responses are synergistic and interactive in response to a polymicrobial infection, which is distinct from responses to a mono-
Bacterial content from oral swabs.

| Pathogen   | Control | Infection |
|------------|---------|-----------|
| P. gingivalis | 45 (100%) | 299.89 ± 100.69 |
| T. denticola | 6 (100%)  | 191.06 ± 68.67 |
| T. forsythia | 6 (100%) | 0.00 ± 0.00 |
| F. nucleatum | 6 (100%) | 0.00 ± 0.00 |

### Table 1. Bacterial content from oral swabs.

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A polymicrobial infection model more closely approximates the natural pathogenesis of the disease. However, prior polymicrobial mouse models of periodontal disease used genetically and immune compromised/modified mouse strains. In an effort to develop a mouse model that more closely resembles the pathogenesis of periodontal disease, this study employed a common mouse strain (BALB/cByJ) with an intact immune system and not genetically altered, to induce periodontal disease with four key periodontal pathogens. Similar to other mouse models of periodontal disease, this current model exhibits the classic signs of alveolar bone loss. The current model further shows the presence of intrabony defects.

Mouse models of periodontal disease have shown various levels of bacterial colonization efficiency similar to the present study. The current study revealed that all four bacterial pathogens (P. gingivalis, T. denticola, T. forsythia, F. nucleatum) colonize the oral cavity of the infected mice based on the PCR data, although T. denticola is less effective. Also, given that the four pathogens are detected at different levels in the oral cavity and in different numbers of mice, this indicates a variable colonization efficiency among the different bacteria. For example, oral swab data (Fig. 2) show that P. gingivalis, T. forsythia, and F. nucleatum were detected throughout the experimental period, whereas T. denticola was not detected at 4 weeks, and when detected, it was not always present in all mice. T. denticola was present in low levels throughout the study; consistent with findings in human periodontal disease. Parenthetically, our results (Fig. 2) show that the four human periodontal pathogens were detected in oral swabs at baseline in uninfected mice (Baseline 1). There may be a few reasons for this finding. Other studies have not detected human oral periodontal pathogens in uninfected control mice, although these other studies used regular PCR methods, whereas the current study used the more sensitive quantitative real-time PCR (qRT-PCR) method of detection. Thus, we may have detected a false positive. Consistent with this observation is that the serum antibody response to the periodontal pathogens in the control uninfected groups was near zero, similar to other studies. Another potential explanation is that these were not germ free mice and they may reveal a low level of contamination. For this reason, all mice were treated with antibiotics to eliminate bacterial load and carriage of contaminating species prior to the start of the study.

Demonstration of a serum antibody response to the bacterial inoculum supports the utility of a given mouse model of periodontal disease. Other mouse models have shown that mice mount an antibody response to P. gingivalis, T. denticola, T. forsythia, and F. nucleatum. Similarly, our data confirmed that this mouse strain can be used to effectively mount an antibody response to all four periodontal pathogens when induced by oral inoculation. All four periodontal pathogens induce a systemic antibody response significantly higher than control mice. Although, serum antibody levels to the four periodontal pathogens were all significantly increased, serum levels of IgG levels against T. forsythia, and F. nucleatum were even higher than to P. gingivalis and T. denticola. In previous studies in genetically altered mice, the serum IgG levels to these four periodontal pathogens were also differentially elevated. The bacterial colonization efficiency of the four pathogens (i.e., Fig. 2) and the altered immune response of the different mice strains may contribute to differences in serum antibody responses. In addition, the high levels of antibodies to T. denticola may suggest that T. denticola exhibits a high virulence level, which could induce a high immune response even in the presence of low colonization. Another possibility is that our sampling times may have missed a high immune response even in the presence of low colonization. For this reason, all mice were treated with antibiotics to eliminate bacterial load and carriage of contaminating species prior to the start of the study.
Fig. 3  Serum IgG antibody response to *P. gingivalis*, *T. denticola*, *T. forsythia*, and *F. nucleatum* inoculation in Control and Infection groups. Data represent the means ± s.d. from six mice per group. *, the difference in serum IgG antibody levels for (a) *P. gingivalis*, (b) *T. denticola*, (c) *F. nucleatum*, and (d) *T. forsythia* was significant (*p* < 0.001) compared to the Control group.

Table 2. Immune gene profiling of oral tissues following polymicrobial infection in mice.

| Gene     | RefSeq   | Description                                           | Fold Change | p Value |
|----------|----------|-------------------------------------------------------|-------------|---------|
| Scgb3a1  | NM_170727| Secretoglobin gene super family cytokine; HIN-1       | 8.473       | 0.191   |
| Il18     | NM_008360| Interferon gamma-inducing factor                      | 2.287       | 0.069   |
| Csf3     | NM_009971| Granulocyto colony-stimulating factor                 | 1.782       | 0.126   |
| Il9      | NM_008373| T-cell growth factor P40; cytokine P40                | 1.732       | 0.187   |
| Il17b    | NM_019508| Neuronal interleukin-17-releated factor               | 1.647       | 0.264   |
| Il13     | NM_008355| T-cell activation protein P600                        | 1.643       | 0.297   |
| Il3      | NM_010556| Multipotential colony-stimulating factor              | 1.623       | 0.359   |
| Il20     | NM_021380| Four helix bundle cytokine 10; IL-20                  | 1.601       | 0.380   |
| Ifna2    | NM_010503| Interferon alpha family, gene 2                       | 1.592       | 0.345   |
| Il6      | NM_001314054| B-cell hybridoma growth factor; IL-HP-1               | 1.583       | 0.270   |
| Gdf15    | NM_011819| Macrophage inhibiting cytokine-1                      | 1.581       | 0.189   |
| Il10     | NM_010548| Cytokine synthesis inhibitory factor                  | 1.568       | 0.312   |
| Il21     | NM_021782| Interleukin-21 isoform 2 precursor                    | 1.559       | 0.267   |
| Spp1     | NM_009263| Early T-lymphocyte activation 1 protein               | 1.549       | 0.309   |
| Ifna4    | NM_010504| Interferon alpha family, gene 4                       | 1.528       | 0.408   |
| Bmp5     | NM_007555| Bone morphogenetic protein 5 preproprotein           | 1.523       | 0.064   |
| Osm      | NM_001013365| Oncostatin-M precursor                     | 1.517       | 0.314   |
| Ltb      | NM_008518| Tumor necrosis factor ligand 1C                      | –2.336      | 0.106   |
the colonization detection was performed at different time points during the study, the antibody response to T. denticola reflects a high summative response at the end even though organism colonization fluctuated.

Other mouse models of periodontal disease have also shown an altered host immune response to a polymicrobial infection. 

However, these mouse models utilized genetically altered (ApoE or integrin beta 6 null mice) mice and they highlighted a variety of altered immune and cytokine profiles in serum and in local or organ-specific tissues. Although, the current study also highlighted an elevated host immune cytokine response featuring 18 genes that were differentially altered (Table 2), these were generally different from those highlighted in other studies. Unique to this study is that the most differentially elevated genes, included secretoglobin gene super family cytokine (8.473-fold change) and interferon gamma-inducing factor (2.287-fold change), and the most differentially decreased gene was tumor necrosis factor ligand 1C (−2.336-fold change). Although, interferon gamma and tumor necrosis factor family members have been implicated in such models, the secretoglobin gene super family cytokine has not been highlighted previously in any models of periodontal disease or inflammation; thus, warranting further study.

A long-held tenet in the relationship between periodontal disease pathogenesis and bacterial plaque has been the concept of a close or proximal radius of effect or radius of action. This concept maintains that bacterial plaque can induce bone loss or mediate destructive effects on the periodontium only within a radius of 1.5–2.5 mm, but beyond 2.5 mm there is no effect. The current study, which shows a widening of the periodontal ligament space throughout the ligament and at a distance far removed from the site of bone loss and initial site of bacterial infection, supports a new concept—that the bacterially mediated radius of effect on the periodontium is far greater than previously thought, and it effects or alters deeper aspects of the periodontal ligament space. This latter finding may have significant implications for the pathogenesis of periodontal disease and biomechanical properties of the periodontium.

The virome, preferentially characterized using shotgun metagenomic sequencing, includes bacteriophages and viruses; both endogenous retroviruses and eukaryotic. The virome has recently been recognized as an essential part of the human microbiome, containing the most numerous and diverse microbial entities, and playing important roles in nutrient and energy cycling. Interactions between viruses, bacteria, and host cells have been associated with shifts from health to disease. The human virome has been implicated in the pathogenesis of several chronic diseases, including diabetes, inflammatory bowel disease, human immunodeficiency virus infection, and cancer. The current study demonstrates a significant change in viral diversity and specific viruses associated with bone loss in the context of periodontal disease. In our study, periodontal disease is associated with an increase in the content and richness of mostly the retroviruses (Golden hamster intracisternal A-particle H18; Jaagsiekte sheep retrovirus; Mouse Intracisternal A-particle; Porcine type-C oncavirus; Murine leukemia-virus-related retroviruses; Squirrel monkey retrovirus; Murine leukemia virus). Interestingly, endogenous retroviral proteins have been implicated in promoting autoimmune disease, neurodegenerative disorders, chronic inflammation, and cancer in humans. Indeed, human endogenous retroviruses express proteins that can elicit a dysregulated immune response. Eukaryotic viruses, such as herpes virus, may also stimulate endogenous retroviruses. Similar to our findings, an altered gut virome has been associated with the loss of gut mucosal integrity and activation of an innate immune response in the mouse intestine.

A previous study of saliva and pooled dental plaque from a limited number of patients (n = 7) with periodontal disease showed differences in virome composition between healthy/gingivitis/mild periodontitis patients and those with moderate/severe periodontal disease. Another virome study in humans with limited periodontal samples identified a new family of redondoviridae associated with periodontal disease. Taken in aggregate,
Fig. 5  The periodontal ligament (PDL) space in infected mice was significantly wider when compared to the PDL space in controls. a, b The PDL-space was measured as the distance between the tooth and bone in control (a) and infected (b) groups. PDL space is shown as a color map on the root surface of the control (aii) and infected (bii) teeth, and as a pseudo ligament in virtual sections of both control (aiii), and infected (biii) teeth. Color maps are indicative of distance between roots (*) of molars and alveolar bone of the periodontal complexes of control (a) and infected (b) mice. c Distribution of PDL spaces in control (solid) and infected (dotted) mice (i) indicate that PDL-spaces in infected mice have a larger variance (broaden peak) and a widened PDL space compared to lower variance (narrower peak) and uniform PDL space in controls (ii). d Left and right tables demonstrate peak values and peak widths of the PDL space in control (top row) and infected (bottom row) mice. No significant difference in peak values of respective PDL spaces between groups was observed ($p = 0.18$). However, a significant difference between peak widths ($p = 0.007$) indicated that the PDL-space in infected mice on average was significantly widened (>100 µm) compared to that of controls. Data represent mean ± s.d.
Fig. 6  Polymicrobial periodontal disease is associated with increased viral diversity. These panels demonstrate the comparison of infected and control mice based on bacterial and viral content. Panel a shows the read content (reads per million) based on bacterial species and panel b shows the read content (reads per million) based on viral species. Panels c and d show the bacterial and viral diversity, respectively. In panel b, the difference in the infected (INF) group was significant \( p < 0.05 \) compared to the control (CTL) group. Data is represented using boxplots.

Fig. 7  Specific microbial and viral species are associated with bone loss and increased PDL space. The graphs show the top microbial and viral species that correlate with the level of bone loss (a–h). The x-axes show the abundance of microbes/viruses and the y-axes shows the level of bone loss in mm. The graph in panel i shows the virus that is most highly associated with and an increased PDL space.
Shotgun sequencing methods examine the combined host and microbial DNA and do not enrich for bacterial DNA, whereas 16S sequencing methods specifically enrich for bacterial DNA. Thus, differences in ability to detect changes in bacterial diversity and total bacterial load may also stem from the use of different sequencing methods.

In summary, this study introduces a new polymicrobial mouse model of periodontal disease in a common mouse strain with an intact immune background. Further, this model makes use of four key periodontal pathogens that are introduced via oral inoculation; an improvement over mono-infection and ligature models. In addition, this model successfully recapitulates the bone loss and intrabony defects characteristic of periodontal disease. A unique and significant altered virome was associated with bone loss in this model of periodontal disease. Lastly, a notable finding affiliated with this model and via the use of micro-CT analysis is that a polymicrobial oral infection can significantly impact periodontal ligament width/space at significant distances far removed from the site of infection and bone loss. This latter finding and the unique altered virome may have significant implications for the pathogenesis of periodontal disease and biomechanical properties of the periodontium.

### METHODS

**Periodontal bacteria and polymicrobial inoculum**

*P. gingivalis* FDC 381, *T. denticola* ATCC 35405, *T. forsythia* ATCC 43037, and *F. nucleatum* ATCC 10953 were obtained from ATCC (Manassus, VA) and cultured anaerobically at 37°C. *P. gingivalis* and *F. nucleatum* were cultured in Tryptic Soy Broth (Becton Dickinson, Franklin Lakes, NJ) containing 5 mg/ml yeast extract, 0.5 mg/ml l-cysteine hydrochloride, 5 μg/ml hemin, 1 μg/ml menadione and 5% fetal bovine serum (FBS) (Gibco Thermo Fisher Scientific, Waltham, MA) for 3 days. *T. denticola* was cultured in oral treponeme enrichment broth media (Anaerobe systems, Morgan Hill, CA) for 5 days. *T. forsythia* was grown in tryptic soy broth containing 5 mg/ml yeast extract, 0.5 mg/ml l-cysteine hydrochloride, 5 μg/ml hemin, 1 μg/ml menadione, 10 μg/ml menadione, 10 μg/ml N-acetyl-l-ascorbic acid (Sigma-Aldrich, St. Louis, MO), and 5% FBS (Gibco) for 7 days. The concentration of each bacterium was determined quantitatively, and the organism was resuspended in phosphate-buffered saline (PBS) at 1 x 10^10 bacteria per ml for experiments.

For the oral polymicrobial inoculation, *P. gingivalis* was gently mixed with an equal quantity of *T. denticola* and allowed to interact for 5 min. Subsequently, *T. forsythia* was added to the tubes containing *P. gingivalis* and *T. denticola*, and the bacteria were mixed gently for 1 min and allowed to interact for an additional 5 min. Lastly, *F. nucleatum* was added and mixed well with *P. gingivalis*, *T. denticola*, and *T. forsythia*. After 5 min, an equal volume of sterile 4% (w/v) carboxymethyl cellulose (CMC) (Sigma-Aldrich) in PBS was added to the bacterial consortium and mixed thoroughly, then this mixture was used for the oral inoculation.

**Infection of mice**

A total of 2 eight-week-old BALB/cByJ female mice (The Jackson Laboratories, Bar Harbor, ME) were housed in microisolator plastic cages and randomly distributed into two groups (six mice per group); a control and an infection group (Fig. 1). The experimental protocols were approved by the Institutional Animal Care and Use Committee of the University of California, San Francisco (IACUC APPROVAL NUMBER: AN171564-01B). Briefly, all the mice were initially given trimethoprim (0.17 mg per ml) and sulfamethoxazole (0.87 mg per ml) daily for 7 days in the drinking water and then their oral cavity was rinsed with 0.12% chlorhexidine gluconate (Peridex, 3M, Maplewood, MN) mouth rinse to suppress the native oral microbiota. The polymicrobial inoculum (5 x 10^9 combined bacteria per ml; 1 x 10^8 cells in 0.2 ml per mouse; 2.5 x 10^9 *P. gingivalis*, 2.5 x 10^9 *T. denticola*, 2.5 x 10^9 *T. forsythia*, and 2.5 x 10^9 *F. nucleatum*) was administered topically in the morning for 4 consecutive days every week for a total of 8 weeks. A control solution without bacterial pathogens was administered as the control treatment.

Oral microbial swab samples were collected with micro cotton tips at different time points to evaluate microbial infection. Sample collection took place the day before antibiotic treatment, the day before infection.

and at 1, 4, and 8 weeks post infection. The post-infection samples were collected 3 days after each infection cycle (Fig. 1). The samples were collected by swabbing the oral cavity, teeth, and surrounding gingiva of the mice using a sterile micro sized cotton swab, which was then immersed in 10:1 Tris-EDTA buffer immediately and stored at −80°C until further processing for DNA isolation.

After 8 weeks of polymicrobial infection, mice were euthanized and the blood was collected via cardiac puncture for analysis of antibody response to the periodontal pathogens. The maxillae and mandibles were dissected from each mouse for various analyses.

DNA isolation, ethanol precipitation, and real-time PCR to confirm bacterial infection

Oral swabs were used to evaluate and confirm infection in the mice. DNA was isolated and purified from the swabs using the QIAamp® DNA Mini kit (Qiagen, Germantown, MD, USA) following manufacturers protocols. Specifically, the swabs were placed in a 2 ml microcentrifuge tube, then 600 μl of PBS was added to each sample tube. To this, 20 μl of QIAGEN proteinase K and 600 μl of Buffer AL were added to each sample, then samples were mixed immediately by vortexing for 15 s. Subsequently, samples were incubated at 56°C for 10 min and briefly centrifuged. Then, 600 μl of ethanol (96–100%) was added to each sample, and samples were mixed by vortexing and briefly centrifuged. Subsequently, 700 μl of this mixture was added to the QIAamp Mini spin column and centrifuged at 6000 × g for 1 min. The QIAamp Mini spin column was placed in a clean 2 ml collection tube, and the tube containing the filtrate was discarded.

Then the steps above were repeated by applying up to 700 μl of the remaining mixture to the QIAamp Mini spin column. Then 500 μl of Buffer AW1 were added and samples centrifuged at 6000 × g for 1 min. Next the QIAamp Mini spin columns were placed in a clean 2 ml collection tube and the collection tube containing the filtrate was discarded. Then 500 μl of Buffer AW2 were added and samples centrifuged at full speed (20,000 × g) for 3 min. The QIAamp Mini spin column was then placed in a new 2 ml collection tube and the old collection tube with the filtrate was discarded. This was centrifuged at full speed (20,000 × g) for 1 min. The QIAamp Mini spin column was placed in a clean 1.5 ml microcentrifuge tube and the collection tube containing the filtrate was discarded. Lastly, 150 μl of Buffer AE was added, sample incubated at room temperature for 1 min, then centrifuged at 6000 × g for 1 min.

Ethanol precipitation of DNA was then performed to prepare the sample for subsequent RT-PCR. Specifically, the following were added to each sample in sequential order: 1/10 volume of 3 M sodium acetate, pH 5.2, and 2.5 volumes of 100% Ethanol. Then 2 μl of a 15 mg/μl glycogen reagent (Glycobluetm Coprecipitant) was added to each sample as a coprecipitating agent. This was mixed by vortexing and samples frozen overnight at −20°C. Then the sample underwent microcentrifugation at full speed (20,000 × g) at 4°C for 1 h. The supernatant was carefully pipetted off, and 250 μl of 70% ethanol was added and the sample, and the sample centrifuged at full speed (20,000 × g) for 10 min. The pellets were dried at room temperature for 15 min, then 25 μl of Tris-EDTA buffer was added to the samples.

Standard RT-PCR was used to confirm the presence of the periodontal pathogens in the oral swabs. The four periodontal pathogens and total bacteria were quantified by RT-PCR using primers (Invitrogen; Table 4) corresponding to 16S ribosomal RNA. Tenfold serial dilutions of DNA of known concentration were used to construct standard curves for quantification of periodontal pathogens and total bacteria. The amplification was conducted using a QuantStudio 3 Real Time PCR system (Thermo Fisher Scientific) with a final reaction volume of 20 μl that included the PowerUp SYBR Green Master Mix (Thermo Fisher Scientific), DNA (15 ng/μl), and primers. The optimized thermal cycling conditions were as follows: 95°C for 10 min followed by 50 cycles of denaturing at 95°C for 15 s, annealing and extension at 60°C for 1 min. The melt curve protocol followed with 15 s at 95°C, and then 60°C for 1 min and 95°C for 1 s. Data were analyzed using QuantStudio™ Design & Analysis Software v1.4.3 (Thermo).

Serum antibody analysis

Blood was harvested from all mice the day of euthanasia then centrifuged at 200 × g for isolation of serum. Serum was then stored at −80°C for subsequent assessment of a specific host response in the form of immunoglobulins (IgG) against P. gingivalis, T. denticola, T. forsythia, and F. nucleatum using an enzyme-linked immunosorbent assay (ELISA). For positive controls for the ELISA assay, each of these pathogens was grown to a cell density of 7 × 10^9 cells/ml and harvested by centrifugation (7000 × g, 30 min, 4°C). After washing, the cells were treated overnight with 0.5% formalin in buffered saline, then diluted to 0.3 (OD600 nm) in 0.05 M carbonate–bicarbonate buffer and used as the coating antigen. The amounts of specific IgG antibodies present were determined by using a mouse IgG ELISA quantification kit (Bethyl Laboratories, USA). A pilot test was first performed with serum diluted at 1:100 to confirm reactivity with the bacterial antigens. Then, after coating with the formalin-fixed bacterial cells, wells were incubated with diluted mouse serum (1:100) for 1 h at room temperature, washed with PBS containing 0.05% Tween-20 (PBST), and alkaline phosphatase-conjugated goat anti-mouse IgG and the 3,5,5′-termethyl-benzidine chromogenic substrate reagent were added for detection and measurement of antibody response. Samples were assayed in duplicate and the purified mouse IgG was used to establish a standard curve on each plate for specific IgG quantification.

Immune-gene profiling PCR array

Twelve sacrificed mice had their right maxillary gingiva dissected and stored at −80°C, six were control mice and six were infected or experimental mice. Tissue was powdered with a mortar and pestle under continuous liquid nitrogen. Total tissue RNA was isolated (RNaseasy mini Kit; QIAGEN) and reversely transcribed (SuperScript VILO Master Mix; Thermo Fisher). Gene expression was assessed using the Mouse Common Cytokines RT² Profiler PCR Array (QIAGEN) containing primers for 84 genes. Results were expressed as PCR cycles to threshold (Ct) and normalized to 1 control gene, glucuronidase beta (Gusb). Ct values were used for calculations of fold changes in gene expression according to the 2^ΔΔCt method.

Shotgun sequencing and microbiome/virome data production and analyses

The gingiva of 12 mice (6 controls and 6 infected) was collected and used for DNA extraction and isolation using the QIAamp® DNA Mini kit (Qiagen, Germantown, MD, USA) according to manufacturer’s instructions. DNA purity and quantity was deemed suitable and met quality control measures for standard shotgun sequencing by Novogen, Inc. (en.novogene.com). Shotgun metagenome sequencing library preparation has been done by Novogen, Inc. and libraries have been sequenced on Illumina HiSeq4000 machines. FASTQ files were generated from the sequencing machines and used for the analyses of the microbiome and virome as described below.

Processing raw data and data clean up

The following criteria were used for processing the raw data and cleaning. Low quality bases (Q-value ≤ 38) which exceeded a certain threshold (40 bp by default) were trimmed. Reads which contained N nucleotides over a certain threshold (10 bp by default) were trimmed. Reads which overlapped with adapter over a certain threshold (15 bp by default) were trimmed.

Metagenome assembly

We utilized de novo assembly for each sample as follows. Samples passing quality control were assembled initially using SOAPdenovo (http://soap.

Table 4. PCR primers used for confirmation of bacterial infection.

| Bacterium                  | Primer sequence (5′-3′)                        |
|----------------------------|------------------------------------------------|
| Porphyromonas gingivalis   | F: AGT CGA GTT GCA GAC TCC GAT CC             |
|                            | R: AAC CCA CAT CGG TAG TTG CTA ACA G          |
| Fusobacterium nucleatum    | F: AGA GTG TGA TCC TGG CTC AG                 |
|                            | R: GTC ATC GTG CAC ACA GAA TTG CTG            |
| Tannerella forsythia       | F: T CCC AAA GAC GGC GAT ATC A               |
|                            | R: ACG GTC GGC ATG TCA TTG T                |
| Treponema denticola        | F: AGGGATATGCGACCGTAGTACA                    |
|                            | R: TTGGGGAGCTTAAACCCAACA                    |
| Universal (total bacteria) | F: TTTGGTTAAGTCCCG                           |
|                            | R: ATCCCCACCTCCTCCTC                        |
Genomics.org.cn/soapdenovo.html. The scaffolds were cut off at “N” to get fragments without “N”, called Scaftigs. Clean data for all samples were mapped to assembled Scaftigs using SoapAligner (http://soap.genomics.org.cn/soapaligner.html) and unutilized paired-end reads were collected. Mixed assembly was conducted on the unutilized reads with the same assembly parameter. The scaffigs of each sample and mixed assembly, which were less than 500 bp, were trimmed.

Taxonomy annotation

The following taxonomy annotation scheme was used. We aligned unigenes to the NCBI nonredundant database with DIAMOND to taxonomically annotate each metagenomic homolog (MEGAN). According to the abundance table of each taxonomic level, various analyses were performed using custom scripts by R and Python.

For the microbiome/virome analyses, two mice from the infection group were excluded from the analysis due to their lack of enough sequencing reads from microbial species. Species with average reads of less than 1 RPM (read per million) were filtered out. For the bacterial and viral content, the p-values were computed using a two-sample t test assuming equal variance of samples from the two groups. The Shannon diversity index was used to quantify the bacterial and viral diversity. We performed linear regression to quantify the association of bacterial and viral species, immune genes to bone loss and periodontal ligament space parameters, respectively. A log10 scale was used for bacterial and viral counts, while the original scale was used for other quantities. The Benjamini–Hochberg procedure was used for multiple testing correction and the corresponding FDR values are reported.

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AUTHOR CONTRIBUTIONS

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COMPETING INTERESTS

The authors declare no competing interests.
