Periodontal disease is more common in individuals with rheumatoid arthritis (RA) who have detectable anti-citrullinated protein antibodies (ACPAs), implicating oral mucosal inflammation in RA pathogenesis. Here, we performed paired analysis of human and bacterial transcriptomics in longitudinal blood samples from RA patients. We found that patients with RA and periodontal disease experienced repeated oral bacteremias associated with transcriptional signatures of ISG15 and CD48-high S100A2-positive monocytes, recently identified in inflamed RA synovia and blood of those with RA flares. The oral bacteria observed transiently in blood were broadly citrullinated in the mouth, and their in situ citrullinated epitopes were targeted by extensively somatically hypermutated ACPAs encoded by RA blood plasmablasts. Together, these results suggest that (i) periodontal disease results in repeated breaches of the oral mucosa that release citrullinated oral bacteria into circulation, which (ii) activate inflammatory monocyte subsets that are observed in inflamed RA synovia and blood of RA patients with flares and (iii) activate ACPA B cells, thereby promoting affinity maturation and epitope spreading to citrullinated human antigens.

INTRODUCTION

Periodontal disease (PD) is more common in patients with rheumatoid arthritis (RA), particularly RA with anti-citrullinated protein antibodies (ACPA) (1). PD is a common disease that affects up to 47% of the adult population (2) and results in gingival bleeding with translocation of oral bacteria to blood (3). RA patients with ongoing PD have increased disease activity (4–6) and are more likely to have treatment-refractory disease (7), suggesting that PD may trigger systemic inflammatory pathways that are relevant for ongoing joint inflammation. However, the mechanisms by which PD and oral inflammation may contribute to ACPA development and persistence in RA are unclear.

ACPAs recognize an array of human citrullinated proteins, generated through posttranslational modifications of arginine to citrulline by peptidylarginine deiminase enzymes (PADs), including filaggrin, fibrinogen, α-enolase, histones, and vimentin (8), and are clinically useful for the classification of RA (9). In established RA, synovial ACPAs positively correlate with disease activity (10), and RA patients who are seropositive are more likely to have flares after discontinuation of conventional treatments (11–13). ACPA often precede the onset of arthritis by years, suggesting that loss of tolerance to citrullinated human antigens is an early event in RA pathogenesis (8, 14, 15). In addition, before onset of RA, there is an increase in N-linked glycosylation sites in the variable regions of ACPA-expressing B cells (16–18), and these N-linked glycans may enhance binding to bacterial lectins (19, 20). Whereas epitope spreading, development of high-titer ACPAs, and accumulation of N-linked glycans in variable regions of ACPAs are associated with the progression to established RA, the mechanisms underlying the induction and reactivation of ACPA-expressing B cell responses are not well defined.
Here, we report our observation that RA patients with PD experience frequent bouts of oral bacteremias, that is, episodes of increased oral bacterial RNA in their blood. These oral bacteremias coincided with inflammatory monocyte transcriptional signatures. In RA patients with PD, bacteremias were enriched with the same inflammatory monocyte signatures and antibody effector function pathways. This observation prompted an investigation of the antibody response to oral bacteria, and we found that oral bacteria that are repeatedly detected in RA blood are broadly citrullinated in the mouth and recognized by extensively somatic hypermutated ACPAs encoded by RA blood plasmablasts. Our findings indicate that damage of the oral mucosal barrier mediated by PD results in repeated, spontaneous translocation of citrullinated oral bacteria to the blood, which triggers innate and adaptive immune responses in RA associated with systemic disease flares.

RESULTS

Oral mucosal breaks trigger systemic inflammatory responses

To determine the role of the microbiome in RA patients with PD, we performed bulk RNA sequencing (RNA-seq) analysis on blood samples from RA patients with and without PD, obtained by weekly finger sticks over the course of 1 to 4 years (Rockefeller University longitudinal cohort, five patients, mean n = 67 time points per patient) (table S1). RNA-seq transcripts were first aligned to the human genome (hg38) for differential gene expression analysis. The human-depleted reads were then aligned to a microbial metagenomic database [Web of Life (WoL)] (21), followed by an in silico decontamination pipeline (Fig. 1, A and B, and fig. S1) (22). To ascertain the mucosal source of the bacteria in the blood, we inferred the relative abundances of bacteria from three oral mucosal sites (buccal mucosa, supragingival plaque, and tongue dorsum) and five other body sites (stool, vaginal fornix, anterior nares, left and right retroauricular creases) using SourceTracker2 (23) and matched body site samples from the Human Microbiome Project (HMP) (24). Because the HMP dataset did not use metatranscriptomics, we validated the approach by applying this pipeline to a publicly available metatranscriptomic dataset of human stool samples (25). We determined that nearly all predicted body site attributions were stool (fig. S2).

The inferred relative abundances of oral bacteria in blood were higher in the time points from RA patients with PD as compared with RA patients without PD (Fig. 1, C and D, and fig. S3, A to E). Conversely, there were no differences in the inferred relative abundances of bacteria from other body sites (Fig. 1C). The most abundant oral bacteria detected in blood by RNA-seq were Streptococcus species, mirroring the most abundant species detected in oral swabs by 16S sequencing (Fig. 1E and fig. S3F). Together, these data demonstrate that bacteremias from the oral cavity, but not other body sites, were more frequent in the blood of RA patients with PD compared with the blood of RA patients without PD.

We next compared human gene expression with the inferred relative abundances of bacteria from various body habitats. Because there were very few source contributions or variances in the levels of fornix- and stool-derived bacteria, these two sites were excluded (figs. S1 and S4A). We found that the inferred relative abundances of tongue dorsum, buccal mucosa, and supragingival plaque, but not any other body sites, were associated with human differentially expressed genes (DEGs) (Fig. 1F and fig. S4B). Moreover, these DEGs were enriched for overlapping innate immune, defense, and interferon pathways (Fig. 1G). Congruently, we observed a positive correlation between the percentage of monocytes, a key innate immune cell, and the inferred relative abundances of oral bacteria in the blood of RA patients with PD (Fig. 1H). We validated the robustness of these SourceTracker2-inferred body site contribution estimates by comparing each SourceTracker2-inferred body site contribution to matched log ratios of the RA data comprising the microbe counts determined to be differentially abundant [ANCOM-BC (26)] in the respective HMP body sites over all other decontaminated microbial read counts (fig. S4, C and D; see Materials and Methods). Given that these three oral sites are proximal to each other, breaches in the oral mucosa likely result in synchronous entry of bacteria from the oral cavity into the blood, inducing activation of innate immune cells, including monocytes, in the blood.

Given that this observation was correlative, we tested whether oral bacteria directly trigger similar gene expression responses in vitro. Given that both RA and healthy oral flora are composed of mostly Streptococcus species, we incubated pooled oral bacteria isolated directly from the oral cavities of healthy individuals with whole blood from healthy individuals in vitro, and we measured relative expression of genes concomitant with oral bacteremias in RA patients (MARCO, TNFAIP6, ERAP2, and ISG15) (fig. S5, A and B). We detected robust induction of these genes in whole blood after 6 and 20 hours of stimulation (Fig. 1I). Thus, oral bacteria can induce systemic inflammatory responses if oral bacteria translocate to blood as a result of an impaired oral barrier, as seen in PD, and this response is not specific to patients with RA.

The observed response to the oral bacteria was likely mediated by granulocytes and monocytes, because purified granulocytes and monocytes up-regulated expression of TNFAIP6 and ISG15 after 6 hours of stimulation with oral bacteria (Fig. 1J). In addition, monocytes stimulated with oral bacteria up-regulated ISG15 protein expression (Fig. 1K), an effect mediated, in part, by the activation of FcγR2a caused by immunoglobulin G (IgG) coating of oral bacteria (fig. S5C) (27). Together, antibody-coated oral bacteria in vitro induce a similar inflammatory innate immune gene signature as observed in vivo in RA patients, with PD likely causing direct activation of blood innate immune cells by oral bacteria.

Inflammatory synovial monocyte genes are enriched in blood during RA flares and in response to oral bacteremia

We next sought to determine whether the oral bacteremias in RA patients with PD contribute to RA flares. In our prior study of RA flares (28), we identified enrichment of myeloid-related pathways in blood during flares as compared with baseline. Using the same longitudinal cohort from Fig. 1, we compared gene expression during flares to baseline and identified DEGs for flares from RA patients with and without PD (Fig. 2A). We classified genes with consistent fold change direction in both PD-associated and non–PD-associated flares as “common flare genes.” Conversely, we classified genes with discordant fold change direction in PD-associated and non–PD-associated flares as “periodontal disease–specific genes” (Fig. 2B). PD-specific flare genes, but not the common flare genes, were enriched for myeloid leukocyte migration, immune complex clearance, and Ig binding (Fig. 2C). Moreover, only the PD-specific flare genes, and not the common flare genes, were
Fig. 1. Oral mucosal breaks trigger systemic inflammatory responses. (A) Experimental workflow. (B) Inferred relative bacterial abundances from eight HMP body sites (n = 336). (C) Body site–inferred relative abundances for time points from RA patients with and without periodontal disease. (D) Inferred relative oral abundances in blood for one RA donor with and one without periodontal disease. (E) Relative abundances of bacteria genera from oral brushings (left) and blood (right). (F) Log_{10} adjusted P values versus log fold changes of human gene expression relative to bacterial abundances of the three oral body sites. (G) Enriched GO pathways in differentially expressed human genes from (F) (adjusted P values). IFN-γ, interferon-γ. (H) Percentage of monocytes in blood cell counts compared with inferred relative oral abundances of oral bacteria, Pearson’s correlation. (I and J) RT-qPCRs of mRNA of inflammatory genes in (I) whole blood, (J) granulocytes, monocytes, and lymphocytes (n = 4 to 6) stimulated with oral bacteria versus unstimulated control. (K) Flow cytometry data showing proportion of ISG15<sup>+</sup> monocytes in CD14<sup>+</sup> monocytes incubated with PBS, oral bacteria, and anti-FcγR2a or isotype (n = 8). (B, H, and I) Two-tailed Kruskal-Wallis test, Dunnett-corrected multiple comparisons. (C) Mann-Whitney U test. (K) Within-subjects analysis of variance (ANOVA), Tukey-corrected multiple comparisons. Mean ± SD. *P < 0.05, **P < 0.01, ***P < 0.001, and ****P < 0.0001 or exact value shown.
enriched for antibody effector functions against microbes and myeloid immune responses (Fig. 2C). Together, these data suggest that (i) immune responses to repeated oral bacteremias contribute to RA flares; (ii) oral bacteremias promote an antibody effector response in RA patients with PD; and (iii) immune signatures in response to RA flares are variable, potentially due to distinct triggers of flare.

Given the specific enrichment of the myeloid leukocyte pathway during flares in RA patients with PD, we compared the host transcriptomic response to oral bacteremias and flares with a recently published single-cell RNA-seq analysis of RA synovial macrophages (29). We found that oral bacteremias and flares only in RA patients with PD, but not those without PD, were enriched for genes expressed by monocyte subsets analogous to ISG15+HLA-A10, CD48+/SPP1+, and CD48+/S100A12+ inflammatory synovial macrophage subsets (Fig. 2D). Similarly, the gene signature of MerTK-HLA-A15+ synovial macrophages had a high probability of co-occurrence with species that dominate the oral cavity, including Streptococcus spp. (Fig. 2E), based on MMvec (30), suggesting that oral bacteria promote activation of the inflammatory monocyte subsets that are analogous to macrophages in the inflamed RA synovium.

**Highly mutated RA antibodies bind citrullinated human and bacterial antigens in vitro**

We reasoned that if flares in RA patients with PD were associated with antibody effector functions, RA patients might harbor additional more specific responses to oral bacteria relative to individuals without RA. Given that PD is specifically associated with ACPA+ RA (31), we hypothesized that ACPAs may cross-react with oral...
bacteria. We therefore screened monoclonal antibodies (mAbs) derived from human RA blood plasmablasts (tables S2 and S3) against a panel of human proteins and bacterial lysates characteristic of the oral mucosa, as well as other mucosal sites. The RA mAbs that bound the human citrullinated autoantigens (ACPA mAbs) cross-reacted with a broad array of in vitro citrullinated bacterial lysates, but not native bacterial lysates or the PAD enzyme used to citrullinate in vitro (Fig. 3A and fig. S6), indicating that they were also anti-citrullinated bacterial antibodies (ACBAs). RA antibodies without specificity against citrullinated human antigens (unreactive mAbs) showed little reactivity to any bacterial lyse (Fig. 3A and fig. S6). There was no difference in B cell receptor (BCR) characteristics, such as CDR3 length or hydrophobicity (GRAVY score), between ACPA and unreactive mAbs (fig. S7, A and B). In contrast, the ACPA mAbs harbored more somatic hypermutations (SHMs) in the V-region of the heavy and light chain (VH and VL) genes compared with the unreactive mAbs (Fig. 3B and fig. S7C), which had similar percentages of SHM as antibody responses in other clinical scenarios (32, 33). In addition, we investigated the frequency of N-linked glycan motifs in the antigen-binding fragment (Fab) region,

**Fig. 3. RA antibodies from plasma, synovium, and gingiva bind both citrullinated human and oral bacteria antigens. (A)** Heatmap of normalized mAb reactivity to citrullinated (cit) and native (nat) bacterial lysates and human recombinant proteins as measured by ELISA. Bars below indicate the mutation load in the V-region and predicted N-glycan motifs. (B) SHM frequency in the V-region in ACPA and unreactive mAbs from (A). (C) Quantification of predicted N-linked glycan motifs in ACPA mAbs and unreactive mAbs from (A). (D) Percentage inhibition of mAb binding to citrullinated S. parasanguinis and Veillonella parvula bacterial lysates after preincubation with citrullinated or native human proteins. (E) Reactivity of plasma IgG from RA (n = 33) and healthy controls (n = 40) against citrullinated and native bacterial lysates. Mean ± SD. (F) Reactivity of synovial fluid IgG from RA (n = 25) and OA (n = 25) against citrullinated and native bacterial lysates. Mean ± SD. (G) Cyclic citrullinated peptide (CCP) reactivity of IgG and IgA eluted from oral bacteria from RA patients (n = 13) and healthy controls (n = 13). Mean ± SD. For (B), two-tailed Student’s t test was performed. For (C) and (G), two-tailed Mann-Whitney test was performed. For (D to F), two-way ANOVA followed by Dunnett’s multiple comparisons test was performed. *P < 0.05, **P < 0.01, ***P < 0.001, and ****P < 0.0001.
characterized by the consensus sequence N-X-S/T (34), which are potential sites for glycosylation that facilitate binding to bacterial lectins (19). ACPA mAbs contained increased numbers of predicted N-linked glycan motifs (Fig. 3C and fig. S7D), consistent with prior reports (16, 35).

To confirm cross-reactivity of ACPA mAbs between citrullinated oral bacterial and citrullinated human antigens, we preincubated mAbs with either a citrullinated or native human antigen and then tested reactivity of the mAbs to citrullinated oral bacterial lysates. The citrullinated, but not the native, forms of human enolase and human vimentin inhibited binding to the citrullinated bacterial lysates (Fig. 3D and fig. S8, A and B). Together, these data demonstrate that antibodies encoded by ACPA-expressing plasmas blasts cross-react against both in vitro citrullinated oral bacterial and human antigens.

The data presented above represent studies of individual B cell clones. To determine whether ACPAs in plasma and synovial fluid of RA patients cross-react to citrullinated bacterial antigens, we compared the binding of IgG antibodies derived from plasma \((n = 37 \text{ RA versus } n = 40 \text{ healthy; table S4) and synovial fluid} \ [n = 25 \text{ RA versus } 25 \text{ osteoarthritis (OA)}] \) with native and in vitro citrullinated streptococcal species (Fig. 3, E and F). IgG isolated from RA plasma, but not healthy donor plasma, bound citrullinated but not native *Streptococcus oralis* and *Streptococcus parasanguinis* (Fig. 3E). Similarly, RA synovial IgG, but not OA synovial fluid IgG, bound citrullinated but not native *Streptococcus gordoni* (S. oralis), and *Streptococcus salivar*us (Fig. 3F). We conclude that RA patients harbor antibodies in both their plasma and synovial fluid that cross-react with in vitro citrullinated oral bacterial and human antigens.

To determine whether oral bacteria are targeted by ACPAs in vivo, we tested whether antibodies bound to oral bacteria, when eluted, were reactive for cyclic citrullinated peptide (CCP) using a clinical anti-CCP antibody assay (36). Antibodies bound to oral bacteria in RA patients had higher CCP reactivity in comparison with those in healthy donors (Fig. 3G), demonstrating that ACPAs are bound to oral bacteria in RA patients. Collectively, these data show that ACPAs cross-react with citrullinated human and oral bacterial antigens.

**Oral bacteria are highly citrullinated in vivo**

We next tested to what extent bacteria from various mucosal sites are citrullinated in vivo. Citrullination of bacteria from three sites (oral, fecal, and vaginal) was analyzed by flow cytometry using an anti-citrullinated peptide antibody (Fig. 4A). The proportion of citrullinated bacteria was highest in oral bacteria as compared with fecal and vaginal bacteria (Fig. 4, A and B).

*Porphyromonas gingivalis* encodes bacterial PAD and is enriched in the gingiva of patients with PD (37, 38), whereas *Aggregatibacter actinomycetemcomitans* expresses leukotoxin A, which induces hypercitrullination of human proteins by forming pores in neutrophils (39). To determine whether specific bacteria are preferentially citrullinated, citrullinated bacteria and unbound bacteria were sorted, followed by 16S ribosomal RNA (rRNA) amplicon sequencing. No significant bacterial family enrichment was identified between the two fractions, suggesting broad and indiscriminate citrullination (Fig. 4, C and D, and fig. S9). The most abundant family of oral bacteria in the citrullinated fraction was the Streptococcaceae family, which does not express a PAD ortholog (40), suggesting that citrullination occurred from an exogenous source of PAD.
maturation and epitope spreading, to bind citrullinated human antigens.

The most frequently detected citrullinated bacterial peptide in saliva in vivo was derived from *S. parasanguinis* amylase binding protein A (AbpA), which promotes bacterial binding to the tooth surface (44). To assess the generalizability of the citrullinated AbpA peptide as an antigentic target in other patients with RA, we compared IgG binding of plasma from RA (*n* = 46) and healthy patients (*n* = 89), as well as IgG binding of synovial fluid from RA (*n* = 31) and OA patients (*n* = 34) to citrullinated and native AbpA peptide. Plasma and synovial fluid from RA patients exhibited increased reactivity to citrullinated AbpA peptide when compared with those from non-RA controls (Fig. 5F). In addition, RA plasma absorbed against these three citrullinated bacterial peptides (citrullinated AbpA, citrullinated DnaK peptide, and citrullinated hypothetical protein) showed decreased reactivity to CCP, citrullinated vimentin, citrullinated enolase, and citrullinated histone 2A, demonstrating cross-reactivity (fig. S12). Furthermore, we measured the citrullinated AbpA antibody levels in healthy individuals with no or mild PD, patients with moderate to severe PD, and RA patients with moderate to severe PD. We found that RA patients with PD, but not those with only PD or healthy controls, harbored anti-citrullinated AbpA antibodies (Fig. 5G). Collectively, these data indicate that RA patients develop antibodies that bind citrullinated oral commensal bacterial proteins that are cross-reactive against known human citrullinated autoantigens.

**DISCUSSION**

Our data suggest that PD-associated oral mucosal breaks result in oral bacteremias that trigger innate and adaptive immune responses, which, when repeated over time in a susceptible host, likely contribute to the pathogenesis of RA. Oral bacteremias trigger activation of innate immune pathways, including inflammatory ISG15+ HLADR+ monocytes, that are present in inflamed joints and the blood of patients with PD and RA flare. Whereas the innate interferon response to oral bacteremia is not specific to RA, antibodies to citrullinated oral bacteria (ACBA) are specific to RA. PD affects 47% of the population, whereas RA only affects 1%, indicating that additional host susceptibility factors must play a role (2, 45). Our finding that some RA patients harbor B cells that bind citrullinated bacterial antigens and subsequently undergo affinity maturation and epitope spreading to bind citrullinated human antigen provides insight into the long-known association between PD and seropositive RA. Overall, the results presented here are consistent with a model where PD enables bacterial translocation that, in turn, stimulates innate and adaptive immune responses that contribute to RA pathogenesis.
Fig. 5. Identification of citrullinated bacterial epitopes targeted by ACPA. (A) Mass spectrometry spectra and retention times of citrullinated and native S. parasanguinis amylase peptide from saliva. (B) Reactivity of RA mAbs to citrullinated and native bacterial peptides determined by ELISA. (C) Binding kinetics of RA35 and GLRA35 to citrullinated DnaK (87–106) and citrullinated fibrinogen (556–575). (D) $K_d$ (dissociation constant) values derived from biolayer interferometry of germline (GL) and ACPA mAbs to citrullinated bacterial and human peptides. (E) Lineage tree analysis of ACPA clonal family as predicted by IgTree. $K_d$ values derived from biolayer interferometry of corresponding mAbs to citrullinated bacterial and human peptides. (F and G) IgG reactivity of (F, left) plasma from RA patients (n = 46) and healthy patients (n = 89), (F, right) synovial fluid from RA patients (n = 31) and OA patients (n = 34) (right), and (G) plasma from individuals with no or mild periodontal disease (HT, n = 20), patients with moderate to severe periodontal disease (PD, n = 60), and RA patients with moderate to severe periodontal disease (RA PD, n = 13) against citrullinated and native AbpA (124–144) by ELISA. Mean ± SD. *P < 0.05, **P < 0.01, and ****P < 0.0001. For (F), two-way ANOVA followed by Dunnett's multiple comparisons test was performed. For (G), Kruskal-Wallis test followed by Dunnnett's multiple comparisons test was performed.
Plasmablasts are a relatively rare B cell population in blood, and antigen-specific plasmablasts typically peak 6 or 7 days after vaccination and then drop to baseline within 2 to 3 weeks (46, 47). We previously observed shared clonal lineages of IgA+ and IgG+ ACPA-expressing plasmablasts in RA blood for up to 12 months, far beyond the established duration of an acute immune response (48). Although we have specifically focused on IgG effector functions, given the numerous studies that implicate Fcγ receptor activation in RA (49, 50), the presence of IgA isotype in the ACPA-expressing clonal families is suggestive of a mucosal drive. Nevertheless, it is possible that IgA and IgG ACPAs contribute to inflammation in RA by forming an immune complex with citrullinated bacterial antigens that activate their respective pro-inflammatory Ig receptors.

Our data show that RA ACPA-expressing plasmablasts exhibit extensive SHM that is greater than typical antibody responses to vaccines (32). Furthermore, we demonstrated that this extensive SHM gives rise to ACPAs that cross-bind both citrullinated oral bacterial antigens and citrullinated human antigens. Because B cells accumulate on average one nucleotide mutation per cycle of activation and affinity maturation in the germinal center (51), the extensive mutation burden of ACPA plasmablasts in RA suggests that they may have been repeatedly restimulated by the recurrent mucosal breaches of citrullinated oral bacteria. For the three mAbs that bound the bacterial citrullinated molecular chaperone DnaK (87–106), the inferred germline or ancestral antibodies that bound the bacterial citrullinated molecular chaperone DnaK (87–106) with low affinity had no detectable binding to citrullinated human antigens. Thus, we propose that PD-mediated repeated oral bacteremias stimulate recurrent rounds of ACPA-expressing B cell affinity maturation and SHM, eventually resulting in epitope spread of the ACPA response to target both citrullinated bacterial and human antigen(s).

We additionally found that non–PAD-expressing oral commensal bacteria are broadly citrullinated in vivo in individuals with PD. Culture of neutrophils with non–PAD-expressing Streptococcus species (40) led to bacterial citrullination, indicating that human-derived PADS released from NETosing neutrophils can mediate citrullination of oral bacteria. Our finding that ACPA-expressing plasmablasts target citrullinated commensal oral bacteria that do not necessarily express endogenous PAD(s) suggests that human neutrophil PADS mediate the generation of citrullinated bacterial epitopes, a result that we confirmed with experiments in vitro.

Entry of oral bacteria into blood was associated with innate immune and defense pathways, indicating a subclinical response to oral bacteremias in RA patients with PD. Several groups have demonstrated that oral dysbiosis in RA is characterized by enrichment of Streptococcus species (52) and that specific isolates of S. parasanguinis can induce arthritis in murine models (53). In PD, oral bacteria released into the blood may disseminate to the liver, spleen, joints, and gut (54–56). Congruently, recent studies have reported an enrichment of oral Streptococcus species in the guts of RA patients (57, 58) that correlates with disease activity, further supporting hematogenous transmission of oral bacteria to distant body sites.

The temporal association of oral bacteremia with ISG15+ HLAhigh monocyte gene expression suggests a mechanism by which these monocytes are activated by citrullinated oral bacteria alone or in ACPA-bound immune complexes. In RA patients with PD, the gene signatures of inflammatory S100A12+ and ISG15+ synovial macrophages were enriched during flares in RA patients with PD. Given that these gene signatures partially overlap, the ISG15+ macrophage subset may represent a recently activated monocyte subset that trafficked from blood and further differentiated into a more inflammatory subset in the synovial tissue. The inflammatory monocyte analogous to the ISG15+HLAhigh macrophage was only enriched in the flares of RA patients with PD, indicating that there is variability in the immune signature of flares, potentially due to distinct triggers. Given the heterogeneous nature of RA, oral bacteremias may trigger a subset of RA arthritis flares, particularly for RA patients with PD. Future efforts will focus on elucidating additional mechanisms of flare in patients without oral bacteremia or PD.

There are several limitations of our approach. It was not possible to know whether the bacterial reads that we identified in blood samples were intracellular, extracellular, or exosomal. The majority of the samples used in our study were from individuals with established RA on disease-modifying anti-rheumatic drugs and/or other biologics, and for some RA patients included in this study, their PD statuses were unknown. Further studies involving samples from at-risk individuals and medication-naïve individuals are needed to understand the impact of medication on RA progression related to oral bacterial translocation. In addition, given that ACPAs can be reactive against several citrullinated antigens and a vast number of citrullinated epitopes, we cannot precisely determine the driving epitope for each BCR with absolute certainty. Last, the proteomic analysis likely did not identify all bacterial citrullinated proteins present in the oral cavity given that >95% of proteins were human, and, thus, those identified only represent a fraction of citrullinated bacterial peptides present.

In summary, the findings described here indicate that PD, through repeated mucosal breaks, results in recurring innate and adaptive immune activation that may contribute to the pathogenesis of RA. Our findings suggest that future studies are needed to determine whether improved oral care may provide therapeutic benefit in the management of RA.

MATERIALS AND METHODS

Study design

The objectives of this study were to investigate the relationship between the oral microbiome and RA flares using finger-stick RNA-seq monitoring and B cell repertoire sequencing. The study used cohorts from Rockefeller University, University of Colorado, and Stanford University/VA Palo Alto.

Rockefeller University longitudinal cohort

Five female patients who met the ACR 2010 classification criteria (9) for RA and were seropositive for CCP+ were followed for a minimum of 1 year with weekly finger stick samples and questionnaires to capture RAPID3 scores and changes in medication and dental work (table S1). They were also assessed with monthly clinic visits that included physical exams with assessment of tenderness and swelling in 28 joints. Additional samples were collected during self-reported flare events. This study was approved by the Rockefeller Investigational Review Board (IRB) (#DOR-0833).

Stanford University cohorts

Studies below were approved by the Stanford University IRB (#3780).
Plasmablasts. Plasmablasts were isolated from blood samples collected from 12 donors recruited at the VA Palo Alto who met 1987 American College of Rheumatology (ACR) criteria (59) and who were positive for anti-CCP antibodies (tables S2 and S3).

Plasma. Plasma samples were collected from individuals with RA who met the 1987 ACR criteria (59) and who were positive for anti-CCP antibodies, as well as from healthy donors (table S4).

Synovial fluid. Synovial fluids from 65 RA and OA patients as determined by board-certified rheumatologists were provided by the Stanford Rheumatic Diseases Biorepository.

University of Colorado PD cohort

Plasma samples from RA individuals who met ACR 2010 classification criteria for RA (57) or from normal donors were collected (table S6). The gingival health of all individuals was classified according to Centers for Disease Control and Prevention and American Academy of Periodontology case definitions (60). This study was approved by the Colorado Multiple Institute Review Board (#15-2288).

Clinical phenotype data and RNA-seq of fingerstick blood specimens from RA patients

The five female patients in the Rockefeller University longitudinal cohort self-collected finger stick blood samples. RNA-seq samples (N = 336) (mean n = 67 per patient) were available for the analysis comparing microbial and human gene expression (Fig. 1). RNA was purified using the Globin-Zero Kit (EpiCentre), cDNA libraries were prepared using the Illumina Truseq mRNA Stranded Library Kit, with 11 to 12 polymerase chain reaction (PCR) cycles for 5 to 8 nM input, and sequencing was performed on a HiSeq2500 system (Illumina) or a NovaSeq 6000 (Illumina) with 150–base pair paired-end reads. Transcript abundance was quantitated with salmon using an hg38 ensGene gene model.

Shotgun metatranscriptomic microbial assignments

Host-depleted and quality-controlled output fastq files were then uploaded to Qiita web server (61) (project ID 13456) for per-sample metatranscriptomic microbial classification. Qiita offers a graphical user interface that easily facilitates shotgun metatranscriptomic and/or metagenomic analysis based on Wolka v0.1.1 (62) and its associated “Web of Life” database containing 10,575 microbial genomes of bacteria and archaea (21) through direct genome alignments. A total of 371 blood-derived RA samples were aligned against the WoL reference genome database using SHOGUN v1.0.8 (63), with Bowtie2 v2.4.1 (64) as the back end. This process is equivalent to a Bowtie2 run with the following parameters: --very-sensitive -k 16 --np 1 --mp "1,1" --rdg "0,1" --rfq "0,1" --score-min "L0,0.05." The sequence alignment is treated as a mapping from queries (sequencing data) to subjects (microbial reference genomes). Reads mapped to a microbial reference genome are counted as hits such that the resultant feature table comprises samples (rows) by microbial genome IDs (columns) and concomitant abundances. These microbial genome IDs (“operational genomic units” (OGUs)) provide a shotgun metagenomic equivalent to amplicon sequence variants in 16S rRNA amplicon sequencing data (62). In the case that one sequence is mapped to multiple genomes by Bowtie2 (up to 16), each genome is counted 1/k times, where k is the number of genomes to which this sequence is mapped. The frequencies of individual genomes were then summed after the entire alignment was processed and rounded to the nearest even integer, thereby making the sum of OGU frequencies per sample nearly equal (considering rounding) to the number of aligned sequences in the dataset. The resultant count matrix was saved as a biom file for downstream analyses. In addition, 217 shotgun metagenomic samples from the HMP (65) that were previously used for source tracking against low biomass data (66) were processed through the same Wolka-WoL OGU pipeline as described above.

Microbial source tracking

Using the RA metatranscriptomic and HMP metagenomic data with annotated HMP body site data, microbial source tracking was performed using a Bayesian source tracking model, SourceTracker2 (https://github.com/biota/sourcetracker2) (23). Details of the Bayesian model have been previously described by our laboratory (66). Because metatranscriptomic data were also available for HMP fecal samples (25), we compared whether HMP fecal metatranscriptomic samples would be assigned as similar or equivalent using SourceTracker2 (fig. S2). On the basis of the near-perfect association between HMP metagenomic and metatranscriptomic data (fig. S3C), we proceeded with using the HMP metagenomic data for source tracking with the RA metatranscriptomic data. Using SourceTracker parlance, the HMP samples served as “sources,” whereas the RA samples acted as “sinks,” and the SourceTracker2 algorithm was used to calculate the proportion of each source attributable to each sink. In lay terms, we estimated the proportion of body site data from HMP data attributable to each RA microbiome sample using the Bayesian model. After (i) intersecting the OGUs in our RA microbiome dataset with those in HMP and (ii) converting the data to BIOM table format (67), we applied the SourceTracking2 approach to all RA samples. SourceTracker2 default settings (alpha1 = 0.001, alpha2 = 0.1, beta = 10, restarts = 10, draws_per_restart = 1, burnin = 100, delay = 1) plus a sink rarefaction depth of 9000 were used. The outputs were calculated in terms of mean fractional contributions of each source to each sink.

Differential gene expression and pathway analysis for oral bacteremias

Differential gene expression was performed with limma (68). Genes with low expression were filtered out using the filterByExpr command in the edgeR R package (60). To remove batch effects and unwanted variation due to low RNA yields, the batch and starting RNA amounts were included in the linear model. Normalization factors were calculated using the “TMM” method, and voomWithQualityWeights was used before fitting a linear model with lmFit and eBayes. Genes with an adjusted P value (false discovery rate, using the Benjamini and Hochberg correction) less than 0.01 were considered significant. SourceTracker2-predicted relative contributions or log ratios (described below) from microbial read counts were used as a continuous variable. For Gene Ontology (GO) pathway analyses, goana function in edgeR was used (69). Gene set enrichment analysis was used for interpreting expression profiles (70).

RT-qPCR of stimulated whole blood and isolated immune cell populations

To isolate monocytes and lymphocytes, frozen peripheral blood mononuclear cells (PBMCs) were thawed and washed with 10% fetal bovine serum (FBS) (Corning) in RPMI 1640 (Thermo Scientific).
Fisher Scientific) and resuspended in deoxyribonucleic acid (10% FBS in RPMI 1640 to improve viability. PBMCs were washed and layered over the Percoll gradient (GE) to separate monocytes and lymphocytes. Monocyte-enriched buffy coat was spun down and resuspended in EasySep buffer (STEMCELL Technologies). Monocytes were then isolated using the EasySep Human Monocyte Isolation Kit (STEMCELL Technologies) as per the manufacturer’s instructions. Neutrophils were isolated as described above. Isolated cells were checked by flow cytometry for >95% purity. The oral bacteria were isolated directly from the mouths of healthy individuals or whole blood. All monocytes, lymphocytes, or neutrophils were resuspended at 7 × 10^6 cells per ml and frozen at −80°C before the experiment. For all of the experiments, a pool of oral bacteria from six healthy donors (combined and aliquoted before freezing) was used to stimulate the monocytes or whole blood. All monocytes, lymphocytes, or neutrophils were resuspended at 7 × 10^6 cells per ml and mixed with oral bacteria or medium. After incubation for 6 hours, total RNA was isolated using an RNEasy Plus Mini or Micro Kit according to the manufacturer’s instructions. For experiments with whole blood, whole blood (2.5 ml) was incubated with oral bacteria (~200,000 bacteria cells) isolated from the oral cavity or mock-stimulated for 0.25 to 20 hours with orbital shaking. At designated time points, blood was added to a PAXgene Blood RNA tube (BD Biosciences) and incubated at room temperature for 2 to 24 hours. Total RNA was purified from the whole blood using the PAXgene Blood RNA Kit (Qiagen) as per the manufacturer’s instructions. Total RNA was reverse-transcribed to cDNA using the High-Capacity cDNA Reverse Transcription Kit (Thermo Fisher Scientific). Reverse transcription quantitative PCR (RT-qPCR) was performed using TaqMan Gene Expression Master Mix and predesigned TaqMan qPCR probes (Thermo Fisher Scientific) on a QuantStudio 7 Flex Real-Time PCR Instrument (Thermo Fisher Scientific). Relative mRNA expression was normalized to the levels of glyceraldehyde-3-phosphate dehydrogenase (GAPDH) and expressed relative to the values for control cells using the ΔΔCT method.

**ISG15 expression in blood monocytes**

Blood monocytes were isolated from frozen PBMCs using the EasySep Human Monocyte Isolation Kit (STEMCELL Technologies) as per the manufacturer’s instructions. Oral bacteria were isolated from the oral cavity as described above. Monocultured *S. para-sanguinis* was grown and incubated with neutrophil-conditioned medium with CaCl2 and dithiothreitol (DTT) for 3 hours before producing citrullinated *S. parasanguinis*. ACPA mAbs or isotype control mAb was incubated with citrullinated *S. parasanguinis* before incubation with monocytes. Monocytes were resuspended at a concentration of 4 × 10^6 cells per ml in 10% FBS (Corning), RPMI 1640 (Thermo Fisher Scientific), 25 mM Hepes (VWR), and 1× penicillin-streptomycin, and 100 μl of cells were mixed with 50 μl of oral bacteria or citrullinated *S. parasanguinis*. In certain experiments, anti-CD32A (STEMCELL Technologies) was added. Cells were incubated for 16 to 20 hours. Cells were washed and stained using CD14 allophycocyanin (APC)—Cy7 (BioLegend), Aquazombie (BioLegend), CD45 Brilliant Violet 711 (BioLegend), and ISG15 Alexa 488 (R&D Systems) according to the manufacturer’s instructions. Cells were analyzed on a Fortessa flow cytometer (BD Biosciences). Data analysis was performed using the FlowJo software.

**Cell barcode–enabled sequencing of plasmablasts**

Sequencing of Ig genes from individual plasmablasts was performed using cell barcodes as described previously (48, 71–73). Briefly, CD19'CD3' IgD'CD14'CD20'CD27'CD38' plasmablasts were single-cell–sorted using BD FACSARia3.1 or BD FACSARia3.2 (BD Biosciences) directly into lysis buffer in 96-well plates. RT of RNA with oligo-dT was carried out in separate wells, and unique well-ID barcodes were added via template switching activity of Maxima Reverse Transcriptase (Thermo Fisher Scientific). Barcoded cDNA from each plate was pooled, and heavy chain (IgA, IgG, and IgM) and light chain (kappa and lambda) were amplified in three rounds of PCRs while attaching plate-specific barcodes and sequencing adapters. PCRs were carried out separating heavy chain of IgA, IgG, and IgM as well as kappa and lambda light chain, and separate libraries were generated from each, gel-purified, cleaned with AMPure XP beads (Beckman Coulter), and sequenced on Illumina MiSeq (Illumina) with 2 × 300 paired-end reads.

**Bioinformatic analysis of Ig sequences**

Sequencing data were processed as previously described (48, 71–73). Briefly, FASTQ files were demultiplexed using the MiSeq FASTQ workflow. Poor-quality reads and bases were trimmed, and the paired reads were stitched together. Similar reads sharing the same plate and well IDs were clustered into operational taxonomic units (74). Heavy chain VDJ genes and light Vj chain were aligned to germline sequences using ImMunoGeneTics (IMGT) HighV-Quest (75). V-region SHMs were calculated using IMGT analysis based on comparison with germline V-region genes. SHM frequency was calculated by dividing the number of mutations by the length of the V-region of the corresponding chain. When comparing antigen-specific plasmablast sequences, the SHM frequency full V-region was compared. N-glycan sites in the heavy and light chain variable regions were predicted by NetNGlyc 1.0 (76).

**Generation of recombinant mAbs**

Previously published plasmablast antibody sequences from RA patients (48, 71, 72) were recombinantly produced in human IgG1 backbone to ensure consistency in the characterization assays. In-house production was done as described previously, using the Expi293 Expression System (Thermo Fisher Scientific) with Expi293F cells (48, 71–73). Briefly, constructs including the heavy chain and light chain variable region sequences were synthesized as gBlock gene fragments (IDT) for cloning into pFUSE antibody plasmids (InvivoGen) using the Cold Fusion Cloning Kit (System Biosciences). We used pFUSEss-CHIgHg1 for gamma, pFUSE2ss-CLlg-hk for kappa, and pFUSE2ss-CLlg-hL2 for lambda. The Expi293 Expression System (Thermo Fisher Scientific) was used for transient transfections, and harvested culture supernatants were purified using Pierce Protein A Plus Agarose (Thermo Fisher Scientific).

**Bacterial culture and preparation**

Bacteria strains were grown to mid-logarithmic growth phase (0.4 to 0.9 optical density units), as determined spectrophotometrically at 600 nm, in American Type Culture Collection— recommended media and conditions (table S7). The bacteria were pelleted by spinning at 6000g for 10 min and then washed in phosphate-buffered saline (PBS) before storage at −20°C.
Bacterial flow cytometry and sorting

Oral “brushings” were collected using BBL culture swabs (BD Biosciences) to brush all teeth above the gingiva for about 2 min according to Stanford University IRB (#3780). Oral samples, healthy vaginal fluid (Lee BioSolutions), and healthy fecal swabs (Lee BioSolutions) were stored at −80°C. One oral swab, vaginal swab, or fecal swab was placed in staining buffer, PBS with 1% bovine serum albumin (BSA; Sigma-Aldrich) on ice for 10 min with vortexing. Each sample was spun down (8000g, 10 min, 4°C) and resuspended in 20% normal mouse serum (Jackson ImmunoResearch) in staining buffer for 20 min on ice before labeling with anti-citrulline mAb (clone 1D9, Cayman Chemical). The sample was washed after the primary staining and resuspended in phycoerythrin-conjugated F(ab’)2 donkey anti-mouse (H+L) (Jackson ImmunoResearch) and SYTO BC dye (Thermo Fisher Scientific) for 30 min at 4°C. The bacteria were washed twice and resuspended in 100 μl of staining buffer containing propidium iodide (Sigma-Aldrich) and 6-μm beads (Thermo Fisher Scientific). Bacteria were run on an LSRII or FACSAria cytometer and sorted directly into DNA/RNA Shield (Zymo Research). Samples with low bacteria concentrations were not included in the analysis.

Citrullination of bacteria via neutrophils

Neutrophils were isolated from freshly drawn peripheral blood from healthy donors as previously described (77). Briefly, neutrophils and red blood cells (RBCs) were isolated from PBMC via Ficoll gradient using Ficoll-Paque Plus (GE). The neutrophil and RBC pellet were diluted in 3% dextrans in Hanks’ balanced salt solution (HBBS; Corning). After a 20-min incubation, the neutrophil-rich supernatant was spun down at 3000g, 10 min, 4°C and resuspended in 20% normal mouse serum (BSA; Sigma-Aldrich) on ice for 10 min with vortexing. Each sample was spun down (8000g, 10 min, 4°C) and resuspended in 20% normal mouse serum (Jackson ImmunoResearch) in staining buffer for 20 min on ice before labeling with anti-citrulline mAb (clone 1D9, Cayman Chemical). The sample was washed after the primary staining and resuspended in phycoerythrin-conjugated F(ab’)2 donkey anti-mouse (H+L) (Jackson ImmunoResearch) and SYTO BC dye (Thermo Fisher Scientific) for 30 min at 4°C. The bacteria were washed twice and resuspended in 100 μl of staining buffer containing propidium iodide (Sigma-Aldrich) and 6-μm beads (Thermo Fisher Scientific). Bacteria were run on an LSRII or FACSAria cytometer and sorted directly into DNA/RNA Shield (Zymo Research). Samples with low bacteria concentrations were not included in the analysis.

Bio-layer interferometry and thermostability assay

Antibody binding to peptide was assessed using ForteBio Octet QK. The binding assays were performed at 30°C and with agitation set at 1000 rpm. Biotinylated peptides were loaded on High Precision Streptavidin Biosensors (ForteBio) at 100 nM concentration in 1× kinetic buffer (ForteBio). The sensors were then quenched using biocytin (Thermo Fisher Scientific). Antibody concentrations ranging from 10 to 8000 nM were measured for each peptide.

Recombinant peptide ELISA

For experiments with the biotinylated peptides, Nunc Immobilizer Streptavidin 384 plates (Thermo Fisher Scientific) were loaded with 1 μM of biotinylated peptide (table S8) for 1 hour at room temperature. Plates were blocked with 50% ChonBlock (Chondrex) or 2% BSA in PBS for 1 hour at room temperature. After blocking, plasma or synovial fluid was diluted at 1:100 concentration or to 1 μg/ml for mAbs and was incubated with shaking at room temperature for 1.5 hours. Dilution series of a positive control serum was included on each plate to ensure consistency between plates. Antibody binding was detected using a horseradish peroxidase–conjugated goat anti-human IgG Fc antibody and Super AquaBlue ELISA substrate. Five washes with PBS–Tween 20 were performed between each step. Blank wells were subtracted for analysis.


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