Biomarkers common for inflammatory periodontal disease and depression: A systematic review

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ABSTRACT

Background: Dysregulated immune response arising in the periphery can induce depressive symptoms through neuroimmune interactions. Inflammatory oral pathology can be a potent inducer of chronic neuroimmune response relevant to depression. We aimed to synthesize available evidence for the association between inflammatory periodontal diseases (IPD) and major depression (MD) in relation to a broad range of biomarkers.

Methods: Medline, Embase, PsycInfo, Cochrane Library, Web of Science and Scopus databases were searched from inception until January 27, 2022. Search terms included subject headings and synonyms for inflammatory periodontal disease and depression. Studies that reported data on both depression and inflammatory periodontal disease as categories along with measurement of a biomarker were considered. Two reviewers independently selected the articles for inclusion, extracted data and assessed the quality of each study. The protocol for this study was registered with PROSPERO, CRD42021215524.

Results: Twenty-eight studies were included in the final review—eleven cross-sectional studies, seven case-control studies, and six prospective cohort studies conducted in humans; the remaining four were experimental animal studies. Eighteen studies including all animal studies reported a positive association between depression and periodontal disease; one study reported a negative association and another nine studies found no such associations. Twenty studies reported mixed associations between IPD and biomarkers (i.e., salivary, serum, urine or gingival crevicular fluid cortisol, C reactive protein, cytokines, etc.). Biomarkers related to depression were gingival crevicular fluid cortisol, interleukin 6 (IL-6), IL-1β, immunoglobulin G against Bacterioides forsythus; root canal lipopolysaccharides; blood IL-6, IL-1β, cortisol, advanced oxidation protein products, nitric oxide metabolites, lipid hydroperoxides and trapping antioxidant parameter; whereas five studies found no associations between depression and a biomarker. Although animal studies showed interaction of immune, inflammatory and neurotrophic biomarkers in the relationship between depression and periodontal disease, human studies showed mixed findings. In most studies, there were risks of bias due to the sample selection and assessment protocol. Study heterogeneity and limited number of comparable studies reporting on shared biomarkers precluded a meta-analysis.

Conclusion: Immune-inflammatory contribution to depression was evident in the context of inflammatory periodontal diseases, but whether biomarkers mediate the associations between IPD and MD needs to be tested through methodologically rigorous studies aiming specifically at this hypothesis.

1. Introduction

For a long time, depression and periodontitis remained appraised as diseases localized within the cranial and oral cavities, respectively. However, recent findings suggest that pathological processes involved in these conditions have far-reaching consequences. In particular, the field of psychoneuroimmunology has established that inflammatory signals arising in the periphery can reach brain areas responsible for mood and behavioral regulation (Raison et al., 2006). In this context, oral pathology including periodontal diseases can be a putative source of neuro-immune dysregulation observed in depressive illness (Hashioka et al., 2019). The evidence base for this hypothesis remains to be established.
Considerable amount of data suggests an increased occurrence of mental health issues in persons with inflammatory oral diseases, and vice versa (Aldosari et al., 2020; Choi et al., 2021; Zheng et al., 2021). A 12-year follow up study of adults (N = 12,708) with newly diagnosed periodontitis showed a 73% increased risk of depression during the follow-up period regardless of sex, age and other comorbidities at baseline (Heu et al., 2015). Most studies included in a systematic review (Peruzzo et al., 2007) showed a positive relationship between stress/psychological factors and periodontal diseases. Furthermore, associations between psychological stress and periodontitis are likely related to or potentially mediated through salivary and blood cortisol, lipopolysaccharides and other markers of cellular and systemic stress and inflammation (Gomes et al., 2018; Goyal et al., 2011; Hilgert et al., 2006; Ishisaka et al., 2008). Stress causes immune responses that increase susceptibility to infection and potentially contribute to the progression of periodontitis (Warren et al., 2014). In an animal model of depression, periodontal ligature-induced bone loss was greatly increased while exhibiting psychological, behavioural and neurochemical responses consistent with depression (Breivik et al., 2006). Together, this evidence indicates bidirectional causal pathways for the comorbidity between depression and periodontal disease and for shared inflammatory mediators.

Altered levels of immune-inflammatory markers including pro-inflammatory cytokines, oxidative and nitrosative stress markers, neurotoxic metabolites of tryptophan degradation and reduced neurotrophic levels are detected in a considerable proportion of individuals exhibiting psychological, behavioural and neurochemical responses consistent with depression (Aldosari et al., 2020; Osimo et al., 2019; Raison and Miller, 2011). The proposed sources of inflammation in depression and periodontal diseases (IPD) as a plausible source of such inflammation in depression. Firstly, IPD involves dysbiosis of the oral microbial community in favor of inflammatory peri-

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**Fig. 1.** Schematic showing biomarkers as putative connecting links between inflammatory periodontal disease and depression. Comorbidity between depression and inflammatory periodontal diseases can be explained in terms of psychological, social, pharmacological and biological, including genetic factors. The bidirectional associations are potentially mediated by a range of inflammatory biomarkers such as cytokines, chemokines, cytokotoxic products of metabolic pathways, stress biomarkers, products of oral/gut microbiome, osteoclastogenic factors and epigenetic regulators. Factors responsible for inciting periodontal and depression pathologies may originate from oral microbes, periodontal tissue as well as the central nervous system and systemic circulation. CRP: C-reactive protein, HMGBl: high mobility group box 1, IL: interleukin, TNF-α: tumor necrosis factor alpha, TLR: Toll-like receptor, NF-κB: nuclear factor kappa-light-chain-enhancer of activated B cells, Th1 cells: T Helper 1 Cells, Treg: Regulatory T cells, LPS: lipopolysaccharide, ncRNA: non-codingRNA, LBX2-AS1: LBX2 Antisense RNA 1, ANRIL: Antisense Noncoding RNA in the INK44 Locus.
signaling in depression as well as afferent inflammatory signals arising from IPD point towards a compelling biological pathomechanism for bidirectional associations between depression and IPD (see Fig. 1). It is of great relevance to study whether empirical evidence supports such mechanistic explanations for these frequently co-occurring and burden-some disorders of great societal significance. We are not aware of any other research projects that directly test this plausible but novel hypothesis. Thus, the objective of this study is to summarize the evidence on the association between inflammatory periodontal diseases and depression mediated by or related to a broad range of biomarkers that are detectable in circulatory and local fluids.

2. Methods

2.1. Search strategy and selection criteria

We conducted a systematic review and attempted meta-analysis following the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) (Moher et al., 2015). Records were searched in the databases Medline (Ovid), Embase (Ovid), PsycInfo (Ovid), Cochrane Library, Web of Science and Scopus from database inception to 27. Jan 2022. Search strings included a wide range of subject headings and synonyms for inflammatory periodontal disease (such as gingivitis, necrotizing ulcerative gingivitis, periimplantitis, periodontitis or periodontal pocket, periocoronitis or gingival hemorrhage) and depression (such as depression or depressive disorder). Full details of the search strategy are available in the Supplementary material (Appendix 1.0-1.1). Two authors screened the title and abstract of 1557 reports independently using Rayyan-a web-based screening solution (Ouzzani et al., 2016).

Inclusion criteria: (a) original peer-reviewed studies conducted in vivo, in humans or animal models, (b) either sex, (c) any age, (d) any human or animal samples, (e) examined both inflammatory periodontal disease and depression, and (e) reported on at least one biomarker. A broad definition of biomarkers was used as suggested by the US National Institutes of Health (National Institutes of Health, 2001). Exclusion criteria: (a) qualitative studies, (b) case reports, (c) opinion articles/letters to the editor, (d) conference proceedings/reviews, (e) studies without biomarker levels analysis, (f) studies where average values and spread of the biomarker levels at group or individual levels not presented or unavailable after contacting the corresponding author, (g) full text not accessible, and (h) articles published in languages other than English or a Scandinavian language. For cases where reports were evaluated differently between two authors, a third author was invited to resolve. Fifty-four articles that screened positive were full text reviewed by two authors. If relevant data were missing from a report, the corresponding

![Identification of studies via databases and registers]

Fig. 2. Preferred Reporting Items for Systematic Reviews and Meta-analysis flow diagram for the current study.
authors were contacted and were requested for additional information. We excluded 26 reports due to violation of inclusion criteria (Fig. 2), leaving 28 reports. The review was registered on PROSPERO (CRD42021215524) before the systematic review was done.

2.2. Data extraction and analysis

Data were independently extracted by all authors in pairs using a standardized form. A third reviewer crosschecked the extraction sheet for each study for consistency and any discordance was resolved through discussion among the authors.

We included studies under an umbrella of depression for any categories the authors defined as satisfying depression diagnosis or having adequate symptom load for depression. We operationalized IPD with any criteria for periodontal inflammation presented by the authors, but included studies that reported values of at least one of the three parameters, i.e., bleeding on probing (BOP), pocket probing depth (PPD) and clinical attachment level (CAL) necessary for a periodontal diagnosis.

We closely evaluated the data against study design, study population and settings for possible meta-analysis of biomarkers reported in comparable studies. As a priori, at least three methodologically comparable studies reporting the same biomarkers would provide basis for further meta-analysis.

2.3. Quality assessment

Key confounding variables considered in each study are presented in Supplementary material (Appendix 2). Potential risks of bias in animal studies were assessed using the SYRCLE’s RoB tool (Hooijmans et al., 2014), an adapted version of the Cochrane RoB tool. Observational cohort, cross-sectional and case-control studies on humans were assessed using the Quality Assessment Tools of the National Heart, Lung, and Blood Institute. The guidelines are available through https://www.nhlbi.nih.gov/health-topics/study-quality-assessment-tools, and the tools contain 14 criteria (cohort and cross-sectional) or 12 criteria (case-control studies) assessing potential selection, information and measurement biases or confounding factors, rated as yes, no, not reported, cannot determine, or not applicable. One point was given for yes, and 0 point for the other responses. Two authors graded each study independently, and the grading was subsequently discussed to reach consensus. Risk of bias matrices are presented as supplementary material (Appendix 3a,3b,3c).

3. Results

3.1. General study characteristics

Of the 28 studies included in the present systematic review, eleven were cross-sectional studies, seven were case-control studies, and six were prospective cohort studies conducted in humans. Remaining four studies were conducted in animals. Human studies were mostly based on dental in- or outpatients and the sample size varied between 30 and 600 individuals. Table 1 Shows detailed information about the study design, population characteristics, sample size, periodontal and depression status, diagnostic criteria as well as the association between periodontal disease and depression.

3.2. Association between IPD and MD

All four animal studies included in this review reported a positive association between depression-like behavior and periodontal disease. Among human studies, fourteen studies found a positive association between periodontal disease and depressive disorder or depressive symptoms suggestive of MD diagnosis. Cakmak et al. (2016) reported elevated depression scores in patients with aggressive periodontitis but not in patients presenting with chronic periodontitis. Nascimento et al. (2019) found higher risk of periodontitis in patients with depressive symptoms but not among those with depression diagnosis, while Gomes et al. (2018) reported higher depression scores among patients with chronic apical periodontitis. Fenol et al. (2017) found significant associations between high levels of stress (including depression) and worse periodontal status. Finally, depression in chronic periodontitis was associated with higher periodontal destruction (Zhang et al., 2021) and stress levels indicative of depression were found to be associated with poorer periodontal healing among IPD patients (Petit et al., 2020, 2021).

Nine of the included studies found no associations between depression and periodontal disease. Yet another study (Johannsen et al., 2006) did find an association between depression and dental plaque and gingival inflammation but did not observe any significant associations for other clinical periodontal parameters (BOP, PPD, CAL, number of teeth). Only one study (Rodriguez Franco et al., 2020) found depression to be negatively associated with periodontal disease.

3.3. Biomarkers

Details about measured biomarkers, sample source and assay method are presented in Table 2. The studies varied greatly between biomarkers that were investigated, type of sample that was used and assay method. The biomarkers reported in the studies generally fell into one or more of the following categories: markers of psychological stress, oxidative and nitrosative stress and damage, bacteria or endotoxin, immunoglobulins, hormones or their receptors, neurotrophic factors and inflammatory cytokines. Cortisol was measured in the majority of the studies, either in saliva, serum or gingival crevicular fluid.

3.4. Biomarkers by IPD and MD

Two studies that used animal models for periodontal disease found elevated inflammatory signaling and reduced neurotrophic support in the brain compared to that of control animals as indicated by higher TNF-α, IL-1β, IL-6 and TLR-4, increased number of activated glial cells as well as reduced BDNF (brain-derived neurotrophic factor) levels (Martínez et al., 2021; Wang et al., 2019). Similar findings were reported for depression-like behavior in animals (Martínez et al., 2021; Wang et al., 2019). Glucocorticoid receptor expression was reported to be decreased in the hippocampus of depression model rats relative to controls (Breivik et al., 2006, 2015).

Twenty of the included studies on human subjects reported on biomarkers related to periodontal disease, fifteen of which found an association between the measured biomarkers and periodontitis. Most studies reported cortisol levels measured in saliva, GCF, serum or urine to be positively correlated with periodontal disease (Bawankar et al., 2018; Cakmak et al., 2014, 2016; Cohen-Cole et al., 1983; Fenol et al., 2017; Rahate et al., 2021; Refullo et al., 2013; Rosania et al., 2009). However, five studies found no associations between salivary cortisol and periodontal disease (Cakmak et al., 2019; da Silva et al., 2015; Kurer et al., 1995; Petit et al., 2020; Zhang et al., 2021). While Zhang et al. (2021) observed no significant difference in salivary IL-1β, Bawankar et al. (2018) found higher serum and salivary IL-1β levels in periodontal patients.

Studies reporting on the GCF and salivary levels of cortisol, MMP (matrix metalloproteinase)-8, MMP-9 showed mixed associations with depression (Johannsen et al., 2006, 2007). Katuri et al. (2016) reported an increase in the levels of serum cortisol in the depressed group. Solis et al. (2016) reported modestly lower levels of IL-6 and IL-1β in blood, and IL-1β in GCF in patients with depression. A significant association between clinical depression and increased levels of root canal LPS, plasma AOPP (advanced oxidation protein products), NOx (nitric oxide metabolites), LOOH (lipid peroxides), and TRAP (total radical trapping antioxidant parameter) was found (Gomes et al., 2018). Depression was not associated with analyzed biomarkers, including cortisol, in five studies (Kurer et al., 1995; Petit et al., 2020, 2021; Rodriguez Franco et al., 2020; Rosania et al., 2009).
### Table 1

Individual study details along with operationalization for depression and inflammatory periodontal diseases, associations between the two and study quality.

| Author (year), country | Sample size | Population, mean age (SD/range), % female | Comparison groups | Inflammatory periodontal disease diagnosis/criteria | Depression diagnosis/criteria | Direction of IPD-MD association | Association between IPD and MD | Study Quality |
|------------------------|-------------|-------------------------------------------|-------------------|--------------------------------------------------|--------------------------------|-------------------------------|--------------------------------|--------------|
| Animal studies         |             |                                           |                   |                                                  |                                |                               |                                |              |
| Breivik et al. (2006), Norway | 40          | Rats, 13 weeks, 0% female                 | Animals with olfactory bulbectomy vs sham operated | Ligature-induced periodontitis (alveolar bone loss) | Experimental (depression-like behavior induced by olfactory bulbectomy) | +ve                            | Periodontal bone loss was elevated (1.00 ± 0.25 mm) in depressed rats vs (0.90 ± 0.13 mm) sham-operated control rats (p < 0.01), and reversed with tianeptine. | See appendix 3a. |
| Breivik et al. (2015), Norway | 43: 12 controls; 31 cases | Rats, 13 weeks, 56% female | Animals with maternal deprivation vs no maternal deprivation | Ligature-induced periodontitis (alveolar bone loss) | Maternal deprivation-induced anxiety and depression-like behavior | +ve                            | Significantly more severe periodontitis in depressed rats. | See appendix 3a. |
| Martinez et al. (2021), Spain | 47          | Rats, 0% female                          | Animals with induced periodontitis or depressive-like behavior alone or in combination | Rat model of periodontitis induced by oral inoculation of *P. gingivalis* and *F. nucleatum* | Experimental depression model using chronic mild stress | +ve                            | Rats with IPD and depressive-like behavior had significantly higher GI values compared to non-depressed rats. In addition, IPD + MD rats had more alveolar bone loss | See appendix 3a. |
| Wang et al. (2019), China | 18          | Mice, 6 weeks, 100% female               | Animals induced with periodontitis vs those receiving vehicle | Mouse model of periodontitis (alveolar bone loss) induced by oral inoculation of *P. gingivalis* | Behavioral mouse model of depression (forced swim and tail suspension tests) | +ve                            | Periodontal disease induced by *P. gingivalis* or LPS caused depression-like behavior in mice. | See appendix 3a. |

Animal studies: Total studies with positive associations between IPD and MD: 4; negative associations: 0; not shown or unclear: 0

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**Cross-sectional studies (Human)**

| Authors (year), country | Sample size | Population, mean age (SD/range), % female | Comparison groups | Inflammatory periodontal disease diagnosis/criteria | Depression diagnosis/criteria | Direction of IPD-MD association | Association between IPD and MD | Study Quality |
|------------------------|-------------|-------------------------------------------|-------------------|--------------------------------------------------|--------------------------------|-------------------------------|--------------------------------|--------------|
| Cakmak et al. (2014), Turkey | 120: 40 control subjects; 41 localized periodontitis; 39 generalized periodontitis | Dental outpatients, 38.3 (24–63) yrs, 51% women | Patients with chronic localized or generalized periodontitis vs healthy controls | Chronic Periodontitis – localized (patients with <7 sites having PD ≥ 5 mm and CAL≥4 mm) or generalized (patients with ≥7 sites having PD ≥ 5 mm and CAL≥4 mm). | Major depression (BDI- Beck Depression Inventory) | –                             | No difference in depression scores between IPD groups | fair |
| Cakmak et al. (2016), Turkey | 92: 31 control subjects; 61 periodontitis | Dental outpatients, 24–60 yrs, 49% women | Patients with chronic periodontitis and aggressive periodontitis vs healthy controls | Generalized chronic periodontitis (PD ≥ 5 mm; CAL≥5 mm at all quadrants and BOP in ≥50% of the sites. Generalized aggressive periodontitis (severe alveolar bone loss and PPD≥6 mm and CAL≥6 mm on ≥8 teeth) | Major depression (BDI- Beck Depression Inventory) | +ve                           | Depression scores significantly elevated in the aggressive periodontitis group, but not in the chronic periodontitis group | good |
| da Silva et al. (2015), Brazil | 64: 21 healthy controls; 43 patients with gingivitis | School students, 11.4 (11–12) yrs., 67% women | Patients with gingivitis vs healthy controls | Gingivitis (based on interview, BOP, Community periodontal index with bleeding on six index teeth) | Depression (CDI- Children’s depression inventory) | –                             | No correlation between MD and IPD. Mean depression score in gingivitis 12.3 (SD 9.1) vs no gingivitis 10.0 (SD 8.6). | fair |
| Fenol et al. (2017), India | 70          | Prison inmates, 38.6 ± 10.9 (25–60) yrs., 0% women | Patients with moderate and severe periodontitis vs healthy controls | Periodontitis Group A (IPD≥4 and 6 mm), Group B (IPD≥6 mm in at least 4 sites) and controls (PPD≤3 mm) | Depression (DASS- Depression, anxiety and Stress Scale) | +ve                           | Significant associations between high stress levels (including depression symptom load) and periodontitis measures. | fair |
|                     | 47          | Dental outpatients,                      | Patients with chronic apical periodontitis (based | Chronic apical periodontitis | Major depression (Beck Depression | +ve                           | Higher depression scores in chronic | fair |

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Table 1 (continued)

| Author (year), country | Sample size | Population, mean age (SD/range), % female | Comparison groups | Inflammatory periodontal disease diagnosis/criteria | Depression diagnosis/criteria | Direction of IPD-MD association | Association between IPD and MD | Study Quality |
|------------------------|-------------|-------------------------------------------|-------------------|-----------------------------------------------|-----------------------------|---------------------------------|-------------------------------|-------------|
| Gomes et al. (2018), Brazil | n=43.5 (18+) yrs., 60% women | Periodontitis with and without depression | On clinical and radiographic exam | Inventory and Hamilton Depression Rating Scale | Apical periodontitis group (BDI 13.5 ± 1.9 vs 2.3 ± 3.1); HDRS 11.5 ± 1.0 vs 3.6 ± 1.7 | Poor |
| Katuri et al. (2016), India | 70: 23 with depression; 24 no depression; 23 yoga-practitioners | All patients with chronic periodontitis divided in 3 groups: with/without depression and yoga-practicing | Chronic periodontitis (based on PI, PPD 5–8 mm, CAL≥5–8 mm) | Depression (ZDS-Zung’s Self-rating Depression Scale) | No difference between the groups with regards to periodontal parameters. | Poor |
| Rahate et al. (2021), India | 90: 30 healthy controls; 30 periodontitis smokers | All patients with chronic periodontitis and stage III periodontitis patients with or without a history of smoking. | Stage III periodontitis (based on PPD ≥6 mm, CAL ≥5 mm, radiographic bone loss extending beyond middle third of the root) | Mild, moderate or severe depression (Zung’s Self-rating Depression Scale) | Patients with periodontitis reported increased depression compared to controls. | Fair |
| Refolio et al. (2013), Peru | 70: 36 with periodontitis; 34 healthy controls | Patients with periodontitis vs healthy controls | Chronic periodontitis (>3 sites ≥ probing pocket depth ≥4 mm). Localized (>30% of teeth) or generalized (>30% of teeth affected). | Depression (Zung’s Self-rating Depression Scale) | All patients with chronic periodontitis received depression diagnosis. | Poor |
| Rodriguez Franco et al. (2020), Mexico | 61: 35 periodontitis; 10 MD; 16 depressive symptoms | Psychiatric inpatients, 46.4 yrs, 51% women | Patients with periodontitis vs patients with depression | Periodontitis (based on BOP, CAL, PD). | Depression symptoms (BDI) and major depression | Fair |
| Rosania et al. (2009), USA | 45 Dental outpatients, 45–62 yrs, 69% women | Recall patients with periodontitis vs patients with periodontitis and depression | Periodontal disease (based on CAL, recession, PPD with severity grades) | Self-reported depression (CES-D) | Positive correlations between missing teeth and depression (r = 0.54, P < 0.001) | Fair |
| Solis et al. (2016), Brazil | 72: 36 MD patients; 36 healthy controls | Psychiatric outpatients, 18–58 yrs, 82% women | Patients with depression vs healthy controls | Periodontal examination (PI, GI, CAL PD, missing teeth, previously reported in Solis et al. (2014)) | Periodontal clinical parameters were indifferent between patients with and without depression. | Good |

Cross-sectional studies (human): Total studies with positive associations between IPD and MD: 6; negative associations: 1; not shown or unclear: 4

Case-control studies (Human)

| Author (year), country | Sample size | Population, mean age (SD/range), % female | Comparison groups | Inflammatory periodontal disease diagnosis/criteria | Depression diagnosis/criteria | Direction of IPD-MD association | Association between IPD and MD | Study Quality |
|------------------------|-------------|-------------------------------------------|-------------------|-----------------------------------------------|-----------------------------|---------------------------------|-------------------------------|-------------|
| Basavakur et al. (2018), India | 75: 25 controls; 50 IPD patients | Healthy controls vs smokers with IPD and non-smokers with IPD | Chronic untreated severe periodontitis (based on CAL ≥5 mm, PPD ≥5 mm, >50% of teeth affected evidence of radiographic bone loss. | Depression (Zung’s self-rating depression score SDS) | Mean depression score significantly higher in IPD than controls. | Fair |
| Cohen-Cole et al. (1983), USA | 70: 35 healthy controls; 35 patients with gingivitis | Patients with gingivitis vs healthy controls | Acute necrotizing ulcerative gingivitis (ANUG, Trench Mouth) (based on pain and interdental papillae necrosis) | Depression (Center for Epidemiologic Studies Depression Scale) | Elevated depression score in ANUG patients. OR for MD in IPD = 4.24 | Fair |
| Johannessen et al. (2006), Sweden | 72: 43 MD patients and 29 controls | Patients with depression vs healthy controls | Clinical periodontal examination (dental plaque, GI, BOP, PPD, CAL, tooth number). Specific criteria not reported. | Major depression (Structured Clinical Interview for DSM-IV) | MD patients had significantly higher amount of dental plaque (0.18 ± 0.13 vs. 0.10 ± 0.10 unit) and higher GI (1.53 ± 0.26 vs. 0.89 ± 0.35) than control subjects. | Fair |

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Table 1 (continued)

| Author (year), country | Sample size | Population, mean age (SD/range), % female | Comparison groups | Inflammatory periodontal disease diagnosis/criteria | Depression diagnosis/criteria | Direction of IPD-MD association | Association between IPD and MD | Study Quality |
|------------------------|-------------|------------------------------------------|------------------|-------------------------------------------------|-------------------------------|--------------------------------|-------------------------------|--------------|
| **Prospective cohort studies (Human)**
| Cakmak et al. (2019), Turkey | 55:15 healthy controls; 40 subjects with periodontitis | Dental outpatients, 40.4 (26-63) yrs., 40% women | Localized or generalized chronic periodontitis (based on PI, GI, BOP, PPD, CAL). Localized: PPD ≥ 4 mm and CAL ≥ 3 mm at ≥ 2 sites and BOP ≥ 20% sites. Generalized: PPD ≥ 5 mm and CAL ≥ 6 mm at multiple sites and BOP ≥ 60% sites. | Major depression (BDI- Beck Depression Inventory) | – | No difference in depression scores after IPD treatment. | fair |
| Kurer et al. (1995), UK | 47 | Dental outpatients, 20-50 (not specified), gender distribution not reported | All patients with high level of oral health | Depression/ anxiety/ psychological mood | – | Depression was associated with plaque level (r = 0.28; p < 0.05), but not gingivitis. | fair |
| Nascimento et al. (2019), Brazil | 539 | Population-based, 31 yrs, 49% women | Prevalence of periodontitis and depression | Major depressive episode (MINI) and depressive symptoms (BDI) | +ve | Higher risk of periodontitis (RR 1.19) and more severe periodontitis in patients with depressive symptoms but not diagnosis. | fair |
| Petit et al. (2020), France | 71 | Dental inpatients, 51.3 yrs, 30-40% women | Patients with depression | Severe generalized chronic periodontitis (based on PI, BOP, PD, CAL with at least 5% of the sites with PD > 5 mm and bone loss). | +ve | Increased stress levels associated positively with worsened outcomes after non-surgical periodontal treatment. | good |
| Petit et al. (2021), France | 71 | Dental outpatients, 51 yrs, 56% women | Periodontitis patients with depression | Severe generalized chronic periodontitis (based on 5% of sites with PD > 5 mm and bone loss). | – | 19% of patients suffered from depression. Increased depression resulted | Good |

Case-control studies (Human): Total studies with positive associations between IPD and MD: 6; negative associations: 0; not shown or unclear: 1.

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Table 1 (continued)

| Author (year), country | Sample size | Population, mean age (SD/range), % female | Comparison groups | Inflammatory periodontal disease diagnosis/criteria | Depression diagnosis/criteria | Direction of IPD-MD association | Association between IPD and MD | Study Quality |
|------------------------|-------------|------------------------------------------|-------------------|--------------------------------------------------|------------------------------|---------------------------------|-------------------------------|--------------|
| Zhang et al. (2021), China | Dental outpatients, 200 IPD only; 200 IPD + MD | Healthy controls versus periodontitis patients with/without depression (smoking). | Radiographic evidence of bone loss. | Chronic periodontitis (based on PPD >5 mm, CAL ≥ 5 mm on more than 30% of teeth and radiographic evidence of bone loss). | Self-reported depression (SCL-90) | – | Depression in chronic periodontitis was associated with higher periodontal destruction, potentially mediated through smoking. | fair |

Prospective cohort studies (Human): Total studies with positive associations between IPD and MD: 2; negative associations: 0; not shown or unclear: 4

Notes: IPD: inflammatory periodontal disease, MD: major depression, BDI: Beck Depression Inventory, HDRS: Hamilton Depression Rating Scale, DASS: Depression, Anxiety and Stress Scale, CAL: clinical attachment level, PPD: probing pocket depth, GCF: gingival crevicular fluid.

3.5. Biomarkers in the relationship between IPD and MD

In animal studies, several of the analyzed biomarkers were related to co-occurrence of periodontitis and depression. Wang et al. (2019) found reduced levels of mature BDNF mediated by increased levels of astrocytic p75NTR in a mouse periodontitis model resulting in depression like behavior (Wang et al., 2019). Breivik et al. (2006) found significantly higher serum levels of cortisol and TGF-1, and decreased TNF-α levels in a rat depression model with ligature-induced periodontitis compared to IPD-only rats (Breivik et al., 2006). Rats presenting both with depression and periodontitis also had significantly higher levels of serum cortisol following LPS stimulation. Furthermore, the same group reported higher hippocampal GR expression in depressed rats with periodontitis, even though there was no association between levels of cortisol, IL-10 or TNF-α and depression (Breivik et al., 2015). Martínez et al. (2021) investigated the expression of a plethora of pro-inflammatory markers including TNF-α, NF-kB, IL-1β, iNOS, mPGES, TLR-4 and markers involved in neuroinflammation (APOA1, corticosterone and GR expression) in the brain, and found several to be dysregulated in depressed/stressed IPD rats compared to controls.

Systemic and urinary cortisol were associated with decreased depression scores in patients with acute necrotizing ulcerative gingivitis (ANUG) (Cohen-Cole et al., 1983). One study concluded that IgG against Bacteroides forsythus (BF) was associated with periodontal disease only in subjects with higher depression scores (Moss et al., 1996). Increased levels of root canal LPS were recorded in patients with chronic apical periodontitis and depression (Gomes et al., 2018). One study reported stress (including depression) and cortisol levels as predictors of attachment loss (Rosania et al., 2009). Patients with both depression and IPD presented with increased levels of cortisol and IL-1β in another study (Zhang et al., 2021). In the study (da Silva et al., 2015), there was a strong correlation between MD and biomarker (diurnal decline in salivary cortisol) in IPD group (r = −0.64; p < 0.01), but not in control group (r = 0.07, NS). Also Rahate et al. (2021) reported on increased levels of salivary cortisol in depressed IPD patients compared to IPD-only patients.

Other studies did not find common biomarkers to explain the link between depression and IPD. Solis et al. (2016) found that the lower levels of cortisol seen in patients with MD were independent of periodontal status. One study concluded that the association between periodontal disease and depressive symptoms is not mediated by systemic inflammation (Nascimento et al., 2019). The activation of proinflammatory immune parameters (IL-1β, IL-6, MMP-8) in periodontal damage was found to be independent of depression in a predictive (hypothetical) model among humans (Rodríguez Franco et al., 2020).

4. Discussion

We examined available literature for the association between depression and periodontitis putatively mediated or related to shared biomarkers. Out of 28 included studies, fourteen human and four animal studies found a positive association between IPD and MD while the rest reported no such associations. A previous systematic review on the relationship between periodontitis and depression selected 15 relevant studies out of which only six concluded on the positive association between these conditions (Araújo et al., 2016), pointing at some progress in this research field. Nevertheless, in our review, we focused on the importance of biomarkers linking the two conditions and not just on their association. Taken separately, most studies found links between periodontal inflammation and cortisol or other biomarker levels, as well as depression and inflammatory markers; yet, in the IPD-MD connection, only ten studies reported on the association of the two conditions via an inflammatory element. Out of these, four studies were performed on animals.

In the current review, animal studies (Breivik et al., 2006, 2015; Martínez et al., 2021; Wang et al., 2019) support the link between depression and periodontal disease via inflammatory biomarkers. This may be due to the ability to subject all study participants to standardized conditions and disease exposures. Nevertheless, there are differences in the presented results. In 2006, Breivik et al. (2006) induced depression by olfactory bullectomy (OB), while in 2015 the same group used the maternal deprivation paradigm for the same purpose. All rats with depression developed a more severe periodontal disease. Yet, in the OB model, rats presented with a decreased glucocorticoid receptor (GR) expression in the hippocampus, while maternally-deprived animals showed increased hippocampal GR expression. This may be indicative of the different degrees of stress and depression that can result in dissimilar biomarker levels that we also see in human subjects. It is considered that an early onset of stressful events, such as in the maternal deprivation model, is thought to consistently change the reactivity to stressors, including immune responses to pathogens (McEwen et al., 2012; Weaver, 2009). This epigenetic modulation of adult human subjects from mixed backgrounds, countries and populations that are included in studies on depression and periodontal disease is difficult to interpret and be adjusted for, contributing to the conflicting results. Based on the available data, it seems plausible that disrupted hypothalamic–pituitary–adrenal axis (HPA axis) and glucocorticoid resistance (i.e., reduced function of glucocorticoid receptor) may directly compromise immune function contributing to the neuroimmune-endocrine pathogenesis of depression (Perrin et al., 2019), as reflected in altered neuroimmune and neurotrophic factors in multiple studies in this review.
### Table 2

Biomarkers reported in each study by depression and inflammatory periodontal disease, and the role of biomarker in the relationship between the two conditions.

| Study | Biomarker | Sample source and assay method | Results: Biomarker by depression | Results: Biomarker by IPD | Role of biomarker in the relationship between IPD and MD |
|-------|-----------|--------------------------------|---------------------------------|--------------------------|-------------------------------------------------|
| **Animal studies** | | | | | |
| Breivik et al. (2006) | Cortisol, TNF-α, TGF-1β, IL-10, GR expression | Serum, hippocampal tissue/RIA (radioimmunoassay) for cortisol; ELISA for TGF-1β; qRT-PCR for GR mRNA | Decreased GR expression in hippocampus of depression model rats. | | IPD + MD rats had significantly higher (1140 ± 388 ng/ml) serum cortisol levels compared to IPD only rats (756 ± 423 ng/ml). IPD + MD rats also had higher TGF-1β (16 ± 4 vs. 12 ± 3) and decreased TNF-α (562 ± 64 vs. 2450 ± 2506) levels. Upon LPS stimulation, compared to controls (627 ± 569 ng/ml), significantly higher serum cortisol levels were found in the depression model rats (1017 ± 606 nm/L), demonstrating that the bullectomy induced a stronger HPA axis responsiveness to the inflammatory LPS. Depression induced hyper-responsiveness of HPA axis (indicated by cortisol level) was not amenable to antidepressant treatment although TGF-1β and TNF-α changes were reversed. Depression models of rat on top of ligature-induced IPD had higher hippocampal GR expression, and lower serum TGF-1β levels after LPS stimulation. TGF-1β levels (pg/ml) in depression group (male: 56 ± 3; female 51 ± 5) vs. healthy control group (male: 63 ± 7, female 58 ± 7); However, cortisol, IL-10 and TNF-α levels did not differ by depression status. Rats with IPD and depression-like behavior had increased expression of pro-inflammatory markers (TNF-α, IL-1β) in the brain. In addition, F. nucleatum was found in the brain parenchyma. These rats also had increased levels of plasma corticosterone and expression of glucocorticoid brain receptors. |
| Breivik et al. (2015) | Cortisol, TGF-1β, IL-10, TNF-α, GR expression | Serum, hippocampal tissue/RIA (radioimmunoassay) for cortisol; ELISA for TGF-1β; qRT-PCR for GR mRNA | Decreased GR expression in hippocampus of depression model rats. | | |
| **Martinez et al. (2021)** | Corticosterone, GR receptor, LPS and 16s rRNA genes of P. gingivalis, P. nucleatum, LPS binding protein, TNF-α, IL-1β, NF-kB (p65 subunit), TLR-4, iNOS, mPGES, phosphor p38 MAPKa/b subunit, APO-A1 phospho-mTOR/mTOR ratio | Plasma, frontal cortex (nuclear extract or homogenate)/ELISA, RT-PCR, Western blot | Rats with depression like behavior had significantly upregulated expression of pro-inflammatory mediators (TNF-α, IL-1β, TLR-4, iNOS and p-p38) in the brain compared to controls. | Rats with IPD had significantly upregulated mRNA expression of TNF-α, and microsomal prostaglandin E synthase (mPGES) compared to controls. | |
| Wang et al. (2019) | Cortisol, p75NTR, BDNF, TNF-α, IL-6, IL-1α | Serum, hippocampus, astrocytes, blood, | Depression-like behavior in periodontitis mice models induced with P. gingivalis had increased number of activated astrocyte and reduced levels of mature BDNF. These effects were reversed by TLR-4 inhibitor TAK242. | P. gingivalis inoculation and LPS from P. gingivalis caused increased alveolar bone loss (mandible) in mice. The mice had significantly elevated serum IL-1α and TNF-α and cortisol levels as well as FPC and hippocampal TNF-α, IL-6 and IL-1α expression compared with the control group. | Periodontal mouse model showed downregulated BDNF maturation through astrocytic p75NTR leading to depression like behavior. |
| **Cross-sectional studies (Human)** | | | | | |
| Cakmak et al. (2014) | Cortisol, DHEA | GCF/ELISA | Higher DHEA (pg/ml) levels in local (64 ± 31) as well as generalized chronic periodontitis (78 ± 39) compared to patient controls without IPD (59 ± 23). Cortisol levels did not differ across IPD groups. | | |
| Cakmak et al. (2016) | Cortisol, DHEA, Salivary flow rate | GCF, Saliva/ELISA | GCF cortisol, saliva cortisol, GCF DHEA and saliva DHEA are elevated, both in generalized | | |

(continued on next page)
Table 2 (continued)

| Study | Biomarker | Sample source and assay method | Results: Biomarker by depression | Results: Biomarker by IPD | Role of biomarker in the relationship between IPD and MD |
|-------|-----------|--------------------------------|----------------------------------|--------------------------|----------------------------------------------------------|
| da Silva et al. (2015) | Cortisol | Saliva/Enzyme immunoassay | and localized chronic periodontitis groups compared to periodontally healthy patients. | No significant differences in the diurnal decline of salivary cortisol between IPD patients and controls (0.17 ± 0.09 vs. 0.24 ± 0.21 ug/dl). | There was a strong correlation between MD and biomarker (diurnal decline in salivary cortisol in IPD group (r = −0.64; p < 0.01), but not in control group (r = −0.07, NS). |
| Fensol et al. (2017) | Cortisol | Saliva/RP Elecsys kit | No significant association between salivary cortisol and depression level | Higher salivary cortisol 26 ± 4 (severe IPD), 19 ± 2 (moderate IPD) vs. control group 9 ± 3 units; P = 0.001). Salivary cortisol correlated positively and significantly with CAL and PP | |
| Gomes et al. (2018) | LPS, LOOH, Nox, TRAP, AOPP, PON1 | Root canal tissue (LPS), plasma/ELISA | There were significant correlations between root canal LPS and depression measured with the HDRS (r = −0.8, p < 0.001, n = 47) as well as the BDI scale (r = −0.8, p < 0.001, n = 47). | Clinical depression was significantly associated with increased root canal LPS, plasma AOPP, Nox, LOOH, and TRAP values, while there were no significant effects of –SH groups and PON1 activity. | Patients with IPD + MD had greatly increased root canal LPS level as compared to IPD-MD group. In subjects with IPD, there were significant correlations between root canal LPS and HDRS (r = −0.7, p < 0.001, n = 34) and the BDI (r = −0.7, p < 0.001, n = 34). Association between depression and IPD was attributable, at least in part, to increased root canal LPS levels in IPD patients. |
| Katuri et al. (2016) | Cortisol | Serum | Serum cortisol levels and depression scores were higher in periodontitis patients with stress vs. with those without stress/ yoga practitioners. | There was a positive correlation between salivary cortisol and depression scores in IPD patients (coefficient r = −0.45 (non-smokers) and 0.40 (smokers)). Ghrelin levels were decreased in depressed IPD patients compared to IPD-only patients. | The more severe the periodontitis, the higher the cortisol levels (OR for periodontitis by cortisol levels 4.14 (95% CI 1.43–12.01) |
| Rahate et al. (2021) | Cortisol, Ghrelin | Serum, saliva/ELISA | There was a positive correlation between salivary cortisol and depression scores in IPD patients (coefficient r = −0.45 (non-smokers) and 0.40 (smokers)). Ghrelin levels were decreased in depressed IPD patients compared to IPD-only patients. | Cortisol levels were significantly elevated in IPD patients compared to controls in serum (16.4 ± 8.9 vs. 11.6 ± 5.5 pg/ml; P < 0.0001) and saliva (399.7 ± 107 vs. 22.55 ± 7.0 pg/ml; P < 0.0001). Ghrelin levels were elevated in IPD patients in both serum (650.25 ± 260.86 vs. 547.6 ± 165.6 pg/ml; P > 0.0001) and saliva (892.4 ± 271.7 vs. 787.3 ± 230.5 pg/ml; P > 0.0001). | The activation of pro-inflammatory parameters in periodontal damage was independent of depression in their predictive (hypothetical) models. |
| Refulio et al. (2013) | Cortisol | Saliva/SCL (high sensitivity electrochemiluminescence) | The more severe the periodontitis, the higher the cortisol levels (OR for periodontitis by cortisol levels 4.14 (95% CI 1.43–12.01) | The more severe the periodontitis, the higher the cortisol levels (OR for periodontitis by cortisol levels 4.14 (95% CI 1.43–12.01) | |
| Rodriguez Franco et al. (2020) | IL1β, IL6, MMP-3 (matrix metalloproteinase)8 | Saliva/ELISA | Depressive symptoms unrelated to proinflammatory immune response | Clinical attachment loss in IPD was associated with pro-inflammatory immune response (a composite of IL-1β, IL-6, MMP-8) | The activation of pro-inflammatory parameters in periodontal damage was independent of depression in their predictive (hypothetical) models. |
| Rosania et al. (2009) | Cortisol | Saliva/RIA (radioimmunoassay) | Cortisol levels alone not associated with depression score. | Positive correlation between Cortisol and higher degrees of periodontal disease measures (probing depth, tooth loss, CAL). Stress and cortisol levels were predictors of attachment loss. | Cortisol & depression in a regression model involving stress were significant predictors of clinical attachment loss |
| Solis et al. (2016) | IL6, IL-1β, INF-γ | GCF, whole blood, stimulated WBC/ELISA | Blood IL-6 and IL-1β were modestly lower in MDD patients compared to controls. WBC upon LPS stimulation showed no differences in cytokine | Cytokine differences in depression were independent of periodontal disease, no mediation analysis was available for IPD-MD associations. (continued on next page) | |

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| Study | Biomarker | Sample source and assay method | Results: Biomarker by depression | Results: Biomarker by IPD | Role of biomarker in the relationship between IPD and MD |
|-------|-----------|--------------------------------|---------------------------------|--------------------------|-------------------------------------------------------|
| Bawankar et al. (2018) | Cortisol, IL-1β | Serum, saliva/ELISA | Salivary cortisol significantly higher in IPD group compared to healthy patient controls (417 ± 100 vs 20 ± 4 pg/ml). Higher serum (19 ± 6 vs. 11 ± 3 and salivary 251 ± 81 vs 160 ± 62 pg/ml). IL-1β levels in IPD patients compared to healthy patient group. Serum cortisol levels not different between IPD and no IPD. |  | Elevated urine and serum cortisol and depression score in IPD patients compared to healthy controls, but no analysis of the three factors together. |
| Cohen-Cole et al. (1983) | Cortisol | Serum, urine/not reported | No significant differences between IPD patients and controls on measures of growth hormone, prolactin, or spot urine catecho-lamines. IPD patients had depressed lymphocyte function, polymorphonuclear leukocyte phagocytosis and chemotaxis. |  |  |
| Johannsen et al. (2006) | Cortisol, IL-1β, IL-6, MMP (matrix metalloproteinase)-8, MMP-9 | GCF, saliva/ELISA, RIA (radioimmunoassay) | MD patients had higher GGF cortisol (3.5 ± 3.3 vs 0.3 ± 0.3 nmol/l) and IL-6 (2.03 ± 1.6 vs 0.79 ± 1.8 pg/site) compared to controls (p < 0.05). MD patients had lower MMP-9 (19.4 ± 12.1 vs 30.6 ± 18.5 pg/site) but GCF IL-1β and salivary cortisol not different between groups. |  |  |
| Johannsen et al. (2007) | Cortisol, IL-1β, IL-6, MMP (matrix metalloproteinase)-8, MMP-9 | GCF, saliva/ELISA, RIA (radioimmunoassay) | Higher GCF IL-6 level in MD vs controls (3.8 ± 1.6 pg/site vs 0.79 ± 1.83 pg/site, p < 0.003), but no difference for IL-1β, MMP8, MMP-9 or salivary cortisol. Patients with MD had lower GCF cortisol than controls. |  |  |
| Karimi et al. (2017) | IgA | Saliva | Salivary IgA level was lower in patients with periodontal disease (207.9 ± 57.2) vs controls (312.66 ± 107.3 units) (P = 0.001). |  |  |
| Leira et al. (2019) | CGRP (Calcitonin gene-related peptide), IL-6, IL-10 | Serum/ELISA | IPD (with migraine) was associated with higher serum CGRP levels (19.7 ± 6.5 versus 15.3 ± 6.2 pg/ml, P < 0.0001) and IL-6 (15.1 ± 9.2 versus 9.6 ± 6.3 pg/ml, P < 0.0001), independent of MD. IL-10 did not show a difference. |  |  |
| Moss et al. (1996) | Antibodies (IgG) against Bf, Aa, Pg | Blood/Antibody assay | IgG against Porphyromonas gingivalis and Actinobacillus actinomycetemcomitans were strongly associated with Periodontitis (OR 4.54 (95% CI 2-10) and 5.3 (95%CI 2-12). |  |  |

**Prospective cohort studies (Human)**

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The possibility of inflammation and depression being related was out of the question a few decades ago (Beurel et al., 2020; Dantzer, 2012). The emergence of psychoneuroimmunology (Ader and Cohen, 1975) and the observed dysregulation in the immune system seen in depressed patients indicated the need for a different approach to the condition (Stein et al., 1985). Elevated levels of IL-6, IL-10, TNF-α or CRP have been associated with depression in different studies (Chamberlain et al., 2019; Irwin and Miller, 2007; Köhler et al., 2017).

Nevertheless, the questions raised are still a matter of debate: is the inflammation primary or does depression promote inflammation? One prospective study on young individuals did indicate that depression episodes increased CRP levels and as such may be considered an inflammatory promoter (Copeland et al., 2012). More evidence is suggesting the role of inflammation in subsequent development of depression (Bondy et al., 2021). Still, a range of other comorbid factors are accountable for an increase in systemic inflammation in patients with depression. Those factors include obesity, smoking, immobility, dysbiotic gut, dental cares or sleep disturbance (Berk et al., 2013). Periodontitis stands out as one such potential factors that has received little attention.

Bacteria are the major culprits in the development of periodontitis, but the tissue response to this aggression results in the secretion and interaction of a plethora of inflammatory mediators contributing to the pathogenesis of periodontal disease (Freire et al., 2021; Suárez et al., 2020). Knowing more on how innate and adaptive immune responses function indicates that the condition is not just a localized pathological event. These biomarkers have been related to the systemic impact of periodontitis and it’s possible association with other conditions (Esteves Lima et al., 2013). At the same time, psychological stress and ineffective coping can also influence the onset and progression of many chronic diseases, including periodontitis through immune suppression (Ng and Keung Leung, 2006; Petit et al., 2020; Zhang et al., 2021). We must not forget that health-related behaviors such as oral hygiene, diet and smoking may be modifiable factors affecting periodontitis and its possible association with other chronic conditions.

The Table 2 contains the summary of the biomarkers and their relationship with depression and periodontitis.

### Table 2

| Study                  | Biomarker                  | Sample source and assay method | Results: Biomarker by depression | Results: Biomarker by IPD | Role of biomarker in the relationship between IPD and MD |
|-----------------------|---------------------------|--------------------------------|---------------------------------|----------------------------|--------------------------------------------------------|
| Cakmak et al. (2019)  | Cortisol, DHEA             | GCF/ELISA                       | Higher cortisol (pg/ml) levels in localized (338.2 ± 309) and generalized (388.0 ± 368) chronic periodontitis compared to patient controls (81.4 ± 27) p < 0.001. No difference across groups at 6-month follow up, nor was DHEA levels different between the groups at baseline and follow up. | | Association between depressive symptoms and periodontitis was not mediated by systemic inflammation. |
| Kurer et al. (1995)   | Cortisol                  | saliva/RIA                      | No associations between cortisol level and depression score | No associations between cortisol level and plaque or gingivitis. | | |
| Nascimento et al. (2019) | CRP                      | serum                          | No significant differences between CRP levels of periodontitis patients and healthy controls, as well as between CRP levels of patients with depression versus non-depressed subjects. | | |
| Petit et al. (2020)   | Cortisol, Chromogranin-A  | serum                           | Stable cortisol and chromogranin-A levels at baseline and 6 months of non-surgical periodontal treatment despite improvement in multiple measures of periodontal outcomes. | | |
| Petit et al. (2021)   | Cortisol, Chromogranin-A  | plasma/immunoassay              | No correlation between plasmatic cortisol or chromogranin-A with psychological status | No correlation between plasmatic cortisol or chromogranin-A with SRP outcomes | |
| Zhang et al. (2021)   | Cortisol, Interleukin B    | saliva/ELISA                    | No significant difference in cortisol or IL-1p levels between periodontitis patients and healthy controls. | IPD-patients (smokers) with depression had increased levels of both cortisol and interleukin B compared to IPD-only patients (non-smokers) and healthy controls. | |

Notes: IPD: inflammatory periodontal disease, MD: major depression, BDI: Beck Depression Inventory, HDRS: Hamilton Depression Rating Scale, BDNF: brain derived neurotrophic factor, CRP: C-reactive Protein, DHEA: dehydroepiandrosterone, INOS: inducible NO synthase, GR receptor: glucocorticoid receptor, AOPP: advanced oxidation protein products, NOx: nitric oxide metabolites, LPS: lipopolysaccharide, LOOH: lipid peroxides, MMP: matrix metalloproteinase, –SH: sulphydryl groups, TRAP: total radical trapping antioxidant parameter, PON1: paraoxonase, WBC: white blood cells, CAL: clinical attachment level, PPD: probing pocket depth, GCF: gingival crevicular fluid.
increase inflammatory markers. Recent evidence suggests that depressed individuals not only have increased baseline inflammation but also show an exaggerated inflammatory response to stress and a diminished responsivity to cortisol. The study by Rosania et al. did use a regression model in which stress was the predictor; in that model, cortisol and stress were shown to be predictors of CAL. In the same context, Petit et al. (Petit et al., 2020, 2021) have concluded that stress associated positively with poor outcomes of periodontal treatment. However, cortisol levels could not be associated with depression scores, nor with periodontal status. Nevertheless, the effects of oral hygiene or periodontal treatments may not be reflective in biomarker levels due to the studied populations; recruited subjects in these studies are adherent to a periodontal maintenance program (Petit et al., 2020; Rosania et al., 2009). Similarly, Kurer et al. (1995) only included patients with good oral health and could not associate cortisol levels to depression or periodontal inflammation. Smoking is a major contributor to both immune and comorbid psychopathologies, requiring a close consideration in all biological assessments of IPD-MD links (Moesintaghavi et al., 2017), which was also reflected in some of the included studies that performed separate analyses for smokers and non-smokers (Bawankar et al., 2018; Rahate et al., 2021; Zhang et al., 2021).

Solis et al. (2016) examined IL-6, IL-1β and INF-γ in GCF as well as systemically and could not prove the hypothesis that immunological activation during depressive episodes contributes to increased levels of cytokines. A certain immunosuppression in MD patients was seen, however. One explanation may be that the sample sizes were quite small, and that types of depression may not be independently distinguished.

Several other methodological limitations of these evaluated studies should be named. First, the majority of studies are observational. It is known that these designs come with the great weakness of not being able to protect against confounding. While most of them used statistical techniques to adjust for possible confounders, there may be residual unknown or unrecognized factors that remain in a heterogeneous adult study population (Smith and Ebrahim, 2003). This is a problem for studies investigating exposures such as depression and periodontitis which can be affected by factors such as health service utilization or socioeconomic status. Cohen-Cole et al. (1983) did find an association between ANUG, depression and elevated cortisol levels; however, the studied population was small, men were underrepresented, and was predominantly of low income. Furthermore, the study design was retrospective. While a statistical adjustment for low income is stated, could other factors related to stressful life events influence the disease progression in the analyzed study population? As such, a causality cannot be concluded, even though an association between depression, acute periodontal inflammation and cortisol levels is noted.

Another critical aspect for the analyzed studies is the big age-span of included subjects. Even though the quality of studies increases if a statistical adjustment is included, other possible unknown age-related confounding factors, particularly multimorbidities associated with immune inflammatory alterations, may influence the outcome of an investigation.

The conflicting results between the investigations may also be explained by the differences between the different levels of periodontal disease severity and the high variation of used clinical parameters. It is also the big variability in studied biomarkers along with testing methodology and limited number of studies for shared biomarkers using the same biological sample that hampered the meta-analysis of existing literature.

In conclusion, there is substantial data supporting associations between periodontal disease and depression and inflammatory contribution to each disorder. However, limited evidence is available on the role of biomarkers in the possible causal pathway between depression and periodontal disease. This is generated by the high heterogeneity among the type of study, the included populations, the assay methods, and the evaluated biomarkers. Future randomized control and prospective studies with standardized clinical and biological assessments are required to establish causality between periodontal inflammation and depression and further analyze the role of biomarkers in linking these diseases.

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**Declaration of competing interest**

None.

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**Appendix A. Supplementary data**

Supplementary data to this article can be found online at [https://doi.org/10.1016/j.jbhid.2022.100450](https://doi.org/10.1016/j.jbhid.2022.100450).

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