Polycystic Ovarian Syndrome and Periodontal Disease: A Systematic Review

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ABSTRACT
Innumerable studies have demonstrated the advantage of non-surgical periodontal therapy in not only improving periodontal health but also in improving the glycemic control levels in patients with type II diabetes mellitus. This study had demonstrated the additive benefits of non-surgical periodontal therapy combined with adjuvant laser therapy on reducing the levels of serum pro-inflammatory cytokines in patients with coronary heart disease. Taking a leaf from these studies it could be hypothesized that non-surgical periodontal therapy could play a vital role in improving periodontal health in patients with PCOS by its positive effect on pro-inflammatory events.

Keywords: Ovarian Syndrome; Periodontal Disease; Cerebrovascular

INTRODUCTION
Polycystic ovarian syndrome (PCOS), a hormonal condition is considered to be one of the common reproductive disorders found in 6-10% of the female population [1] mostly affecting women of reproductive age group, with the prevalence ranging from 9.13% to 36% in India [2,3]. The pathophysiology of PCOS involves primary defects in the hypothalamic-pituitary-gonadal axis; which results in disordered secretion of gonadotropin by the hypothalamus leading to raised luteinizing hormone (LH) levels and normal and /or low follicle-stimulating hormone (FSH). The elevated LH levels, in turn, stimulate the ovarian theca interstitial cells to secrete excessive androgen, which in turn can account for the steroidogenic abnormalities [4]. Women with PCOS were found to have excessive androgen secretion which may cause numerous small collections of soft fluid in the ovaries leading to cyst formation and also have abnormal and prolonged menstrual periods. There are various phenotypes characterized by this syndrome and the three common phenotypic features of PCOS are hyperandrogenism, polycystic ovaries, and ovulatory dysfunction [5]. Along with these leading phenotypic features, this condition can also be manifested with metabolic disorders like insulin resistance (found in 60-80% of women with PCOS) [6], obesity, type 2 diabetes mellitus (found in 5.8% of women with PCOS), dyslipidaemia (found in 93.3% of women with a low HDL (<50mg/dL) being the commonest feature seen in 91.7% of PCOS cases) and increased risk factors for both cerebrovascular and cardiovascular disease [7]. In addition to being an endocrinological disorder, PCOS is also considered a multifactorial disorder and other factors like genetic, metabolic and environmental abnormalities adds to its etiology. Hence women with PCOS have been found to have higher levels of depression and overall psychological morbidity and decreased quality of life due to mood disturbances, decreased sexual satisfaction, weight gain, acne vulgaris, and alopecia.

PCOS is associated with low-grade systemic inflammation and is indicated by elevation of multiple markers of inflammation such as C-reactive protein (CRP), proinflammatory cytokines and chemokines including interleukin 18 (IL-18), monocyte chemoattractant protein-1, macrophage inflammatory protein-1, and white blood count. Furthermore, increased oxidative stress and its biomarkers suggest PCOS as an inflammatory disease [8].

It is a deep-rooted fact that periodontitis is a chronic inflammatory disease and it is the inflammation that links periodontitis with various systemic diseases [6]. C-reactive protein is one of the important markers of inflammation, produced under the stimulatory control of proinflammatory cytokines such as IL-6 and TNF-α. Literatures have suggested that chronic infections associated with increasing levels of
reactive oxygen species, myeloperoxidase (MPO), oxidative stress, inflammatory cytokines (such as IL-6 and TNF-α), high-sensitivity C-reactive protein (hsCRP), adhesion molecules and blood lymphocytes and monocytes have a role in the etiology and pathogenesis of PCOS [3]. In PCOS there will be increased levels of androgens and estrogens which in turn affect the local microbiota sub gingivally and invariably acts on the gingival cells and change the effectiveness of the epithelium [3] leading to gingivitis and periodontitis in PCOS affected women. These sequential events of proinflammatory events and hormonal imbalance have been considered as a possible link between Periodontal Disease and Polycystic Ovarian Syndrome.

However, there is still limited information about periodontal health, oral microbiota, and its association with PCOS. The aim of the present study was to explore this association between PCOS and periodontal health by focusing on the oral aspects of PCOS through a systematic review of indexed literature.

MATERIALS AND METHODS

Design

A systematic review was undertaken using objective and transparent methods as per the PRISMA (Preferred Reporting Items for Systematic Reviews and Meta-Analyses) guidelines, to identify, evaluate and summarize all relevant research findings. The protocol for systematic review was registered first with PROSPERO (International Prospective Register of Systematic Review) Register ID CRD42020171430

Eligibility criteria: On applying the PECO analysis to the articles searched, the criteria were set as shown below:

PECO analysis

Population: Polycystic ovarian syndrome women with the reproductive age group of 15-49 years

Exposure: Proinflammatory cytokines, Oxidative stress biomarkers, and Oral microbiological parameters

Comparison: Healthy women with the reproductive age group of 15-49 years

Outcome: Clinical Periodontal parameters like probing depth (PD), plaque index (PI) and bleeding on probing (BOP)

Inclusion and exclusion criteria

Inclusion criteria:
- Women with the reproductive age group of 15-49 years.
- Women with the history of polycystic ovarian syndrome, as defined by Rotterdam’s criteria [5].
- Cross-Sectional and case-control studies are included.
- Studies published in the past 10 years were included (From 2009-2019).
- Studies which was written in English languages were only included

Exclusion criteria:
- Qualitative studies, reviews, expert opinion, systematic reviews, meta-analysis, and case studies/series.
- Publications with no abstract and those which were widely out of the scope of the study were eliminated.
- Studies that required translation to the English Language.

The remaining studies were sorted out on the basis of their title and abstract. Finally, those studies in which the abstract fulfilled all the inclusion criteria were selected for full-text reading. In those cases, in which a study met the eligibility criteria but the information in the abstract was insufficient, full texts of the articles were also obtained. A further literature search was performed based on the references of the selected articles.

Strategy

To identify the pertinent studies, a broad search of the literature was done using PubMed, Google-Scholar, Tripdatabase, and Cochrane from the year 2009-2019 (Table 1). A detailed search strategy was developed for MEDLINE through the use of MeSH terms and was revised for Google Scholar. The first set of terms include ‘Polycystic ovarian syndrome’, ‘Periodontitis’ separated by Boolean operator OR. The second set included the term ‘women with Polycystic Ovarian Syndrome’, ‘periodontal disease’ separated by Boolean operator “AND” and the third set included the term ‘Association of PCOS and periodontitis,’ separated by Boolean operator “AND”. Hand searches of reference lists of included studies were conducted to ensure additional relevant references. Although systematic reviews, qualitative studies were excluded, reference lists were checked to ensure all primary research was located for inclusion. Only full papers written in English were included. Where multiple publications reporting on the same study existed in different databases, data from the study were extracted and reviewed only once. Duplication of the article was identified using the Zotero.

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Table 2: Modified Newcastle Ottawa scale – cross sectional studies.

STUDY

Study selection was conducted by two authors who independently screened titles and abstracts against the inclusion/exclusion criteria and identified relevant papers. Then the same two authors independently reviewed the full-text studies unable to be excluded by title and abstract alone. A Comparison of papers was completed between the two authors with no disagreements.

DATA EXTRACTION

The data extraction from the final 14 articles was done using a data extraction form. It includes the first author name, year of publication of the article, the aim of the study, objectives of the study, study design, study summary, results, and outcome.

QUALITY ASSESSMENT OF THE INCLUDED STUDIES

The final analysis included 4 cross-sectional studies and 10 case-control studies the methodological quality of the selected articles were assessed using the modified Newcastle Ottawa Scale for cross-sectional studies and the Newcastle Ottawa Scale for case-control studies.

RESULTS

While typing the meSH terms, 1380 relevant articles were identified (PubMed=9, Google Scholar=1360, TRIPDATABASE=7, Cochrane Database=1, EBSCO=3). Thousand two thirty-eight articles were eliminated after reading the title. Four articles were eliminated due to duplication. One forty-two articles were selected for the abstract reading. After the abstract reading, one twenty-one article was excluded and twenty-one articles were included. After reading the full text, seven articles were excluded and fourteen studies that met the inclusion and exclusion criteria were included Figure 1.

Figure 1: Flow diagram of different phases through the systematic review.

Types of participants

The participants included were polycystic ovarian syndrome women with the reproductive age group of 15-49 years. In all studies PCOS was diagnosed according to the criteria of Rotterdam[5] with the presence of at least two of the following: (1) polycystic ovaries (presence of >12 follicles in each ovary measuring 2–9mm in diameter and/or increased ovarian volume>10ml), (2) oligomenorrhea and/or anovulation and (3) hyperandrogenism (Clinical: Acne, Hirsutism, acanthosis nigricans, Biochemical: Total T >70ng/dL, Androstenedione >245ng/dL, DHEA-S>248μg/dL).

Types of outcome measures

Primary Outcome: Clinical Periodontal Parameters. In all 14 studies, a comprehensive clinical periodontal examination was performed using different periodontal parameters like probing depth (PD), plaque index (PI) and bleeding on probing (BOP), Clinical attachment loss (CAL) and gingival index (GI) were reported respectively. One study reported the rate of tooth loss among patients with and without PCOS. Intraoral radiographs were used in four studies and panoramic radiographs were used in one study to assess bone loss. Diagnostic methods and altered clinical parameters of the analyzed studies.

DISCUSSION

All the analyzed studies revealed a definite relationship between PCOS and PD (gingivitis and/or periodontitis), and hence, it is possible to associate both PCOS and PD.

Literature review has demonstrated an increased production of sex steroid hormones like estrogen might lead to increased gingival enlargement, followed by gingival bleeding and microbial changes. Increased estrogen levels could also be noted in the PCOS condition, which could explain its impact on the periodontium as demonstrated in a case report done.
this study, the author compared the gingival biopsy specimen of PCOS patients with and without gingivitis for estrogen and progesterone receptor levels. This study concluded that estrogen receptors were present in the gingival biopsy specimen of PCOS patient with gingivitis with the absence of other receptors in both case and control. Also precisely, estrogen receptors beta (ERbeta) produced estrogenic effects in PDL cells, but there was no immunoreactivity expressions for progesterone receptors in these cells, which implies that progesterone does not have a direct effect on PDL cell function.

Studies have also demonstrated an altered immune-inflammatory response of the periodontium in the presence of various systemic disorders. This altered host response could be attributed to the increased pro-inflammatory cytokines along with hormonal changes leading to an alteration in the oral microflora, eventually, even a sub minimal plaque deposits in a susceptible host could lead to greater periodontal destruction within a shorter period of time. Altered host response in the gingiva may lead to greater periodontal destruction adversely affecting the alveolar bone and adhesive joints, eventually leading to tooth loss. Of late the role of oxidative stress in complementing this disease progress has also been kindled.

Studies have associated the increased prevalence of periodontal disease among patients with certain non-communicable diseases like diabetes and cardiovascular disease. Bleeding on probing could be attributed to the role of oxidative stress on the periodontium. Similar phenomena could also play a vital role in the association between PCOS and PD.

Reported significant positive correlations with all clinical periodontal parameters and oxidative stress markers of MDA levels. Detected that MPO activity was significantly higher in PCOS women which were inconsistent in the study done. Who suggested that MPO levels were not found to be significantly different? The higher BOP rates in PCOS group can be attributed to the influence of hyperandrogenism which causes excessive proliferation of vascular endothelial cells and epithelial keratinization in the gingiva.

Innumerable studies have demonstrated the advantage of non-surgical periodontal therapy in not only improving periodontal health but also in improving the glycemic control levels in patients with type 2 diabetes mellitus. This study had demonstrated the additive benefits of non-surgical periodontal therapy combined with adjuvant laser therapy on reducing the levels of serum pro-inflammatory cytokines in patients with coronary heart disease. Taking a leaf from these studies it could be hypothesized that non-surgical periodontal therapy could play a vital role in improving periodontal health in patients with PCOS by its positive effect on pro-inflammatory events. However, this phenomenon has to be further explored by longitudinal studies by not only assessing subjective oral markers but also an objective clinical assessment of periodontal health.

LIMITATIONS

The outcomes of this systematic study have to be interpreted with the following limitations. Since all studies included in this review were cross-sectional in nature the temporality of this association cannot be well estimated. Being a hormonal disease, PCOS could also initiate a variety of inflammatory biomarkers independent of periodontal disease. Hence, the role of pro-inflammatory markers could always be confounded by the combined effect of disease as both the studies of interest have an inflammatory origin. As a result, a single biomarker assessing the association between PCOS and Periodontal disease could not be demonstrated. The studies included in this review assessed a variety of biomarker making meaningful interpretation difficult. This was a reason why the authors restricted the present study to a systematic review rather than a meta-analysis.

CONCLUSION

Within the constraints of the present design, our systematic review clearly demonstrates an association between PCOS and periodontal disease. All 14 studies included in this review demonstrated this association unanimously. In the absence of a possible explanation for this phenomenon, the authors suggest well designed longitudinal studies which might throw light on the missing link between PCOS and PD.

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