

Is Periodontal Disease a Risk Factor for Coronary Artery Disease (CAD)?

(Les maladies parodontales sont-elles un facteur de risque des coronaropathies?)

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S o m m a i r e

Les coronaropathies (CP) demeurent la principale cause de décès dans la plupart des pays industrialisés, malgré les progrès sensibles qui ont été réalisés en matière de prévention et de traitement. Et les données épidémiologiques actuelles laissent croire qu'on pourra difficilement maintenir les gains récents réalisés dans la réduction de la prévalence de ces maladies, tant que les populations à haut risque ne seront pas mieux ciblées. Bien que les infections dentaires (et principalement les infections parodontales) aient récemment été définies comme un facteur de risque indépendant des CP, les données actuelles sont insuffisantes pour justifier le traitement de ces infections comme moyen de freiner une CP ou autre affection systémique (p. ex., diabète, accident vasculaire cérébral ou issues défavorables de la grossesse) ou d'en renverser la progression.

Mots clés MeSH : coronary disease/etiology; periodontal diseases/complications; risk factors

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A high concentration of plasma cholesterol, especially low-density lipoprotein cholesterol, is the principal established risk factor for the multifactorial pathogenesis of atherosclerosis.¹ Coronary artery (ischemic) disease (CAD) is unquestionably the most important manifestation of atherosclerosis, although the pharmacologic reduction of plasma cholesterol levels and other lifestyle changes have contributed to reductions in age-adjusted mortality rates of only about 50%.² This implies that the pathogenesis of coronary artery atherosclerosis is linked to other risk factors, although the precise identification of these factors is hampered by complex interactions between focal lipid accumulations in arterial walls and associated host responses.¹

More precise identification of these risk factors is further constrained by the lifelong origin of CAD. For example, CAD may initially present as "fatty streak" lesions in the arterial walls of children and adolescents, where the purely inflammatory host reactions result in focal collections of monocyte-derived macrophages and T lymphocytes.³ Hypercholesterolemia in adults is more commonly linked to progressive extracellular deposition of amorphous and membranous lipids

before the influx of inflammatory cells into these atherosclerotic lesions.⁴ Significant declines in CAD mortality rates associated with these atherosclerotic lesions may be attributed to therapeutic reductions in cholesterol levels, but the recent plateau in these rates has been associated with slight resurgences in the prevalence of stroke (cerebrovascular diseases).⁵ The impact of these resurgences may, in turn, exacerbate the difficulties in predicting who will experience the symptoms of CAD. For example, such symptoms tend to be prevalent in only a fraction of those with one or more established risk factors, whereas others succumb to CAD without any apparent risk factors.²

Current research indicates that varied clusters of interactive risk factors contribute to the onset of CAD, thereby complicating the precise assessment of high-risk individuals.⁶ Recent evidence for microbial (notably herpes virus and *Chlamydia pneumoniae*) infections as independent risk factors for CAD clearly illustrates these concerns.⁷ For example, administration of antibiotics significantly reduced the prevalence of ischemic attacks in a recent randomized, double-blind, placebo-controlled trial,⁷ whereas interactions between infections and

other, possibly more important, risk factors may predispose genetically more susceptible individuals to CAD. As a result, current evidence cannot be used to justify the routine prescription of antibiotics for prevention or treatment of CAD.⁸ This review evaluates whether similar reservations apply to the treatment of oral (notably periodontal) infections for the arrest or reversal of CAD.

Is There Evidence for an Association Between Periodontitis and CAD?

Recently published studies that have purportedly designated dental infections as independent risk factors for CAD are difficult to validate. The inherent deficiencies include the following:

- difficulties in the design of studies to accommodate potential interactions between variable clusters of risk factors and other factors that are more ill defined (e.g., physical and mental stress, differential circadian periodicities);

- difficulties in the reliable acquisition of standardized data for dental (especially periodontal) infections for baseline and sequential analysis.

These deficiencies have yet to be comprehensively addressed in published studies. For example, not only do the available studies range from cross-sectional to case-control and prospective designs, but quantitative assessments of dental infections extend from periodontal probing to non-validated indices and self-reporting. Most of these studies suggest that CAD is linked to periodontal infections (Tables 1–3), although inconsistencies in design preclude rigorous assessment (e.g., through a Cochrane review).

Is the Periodontitis-Associated Risk of CAD Reversible?

If the link between periodontal infections and CAD is confirmed by subsequent studies, the question will still remain whether the benefits of treating the infections are analogous to

Table 1 Summaries of case-control studies on the relationships between periodontal infections and coronary artery disease (CAD)

| Patient numbers and characteristics | Results | Comments |
|---|--|---|
| 100 consecutive patients ≤ 50 years of age admitted to hospital for MI and 102 ≤ 50 years without MI as controls ⁹ | | |
| Series 1: 40 men with MI, 41 men without MI as controls | Significant oral health differences ($p < 0.001$) between cases and controls | Assessments of relationships between oral infections and MI compromised by unproven reliability of "total dental index" and "pantomographic index" to quantify oral health |
| Series 2: 44 men and 16 women with MI, 44 men and 17 women without MI as controls | Significant oral health differences ($p < 0.001$) between cases and controls | Assessments of relationships between oral infections and MI compromised by unproven reliability of "total dental index" and "Pantomographic Index" to quantify oral health |
| 85 patients with CAD and 46 random controls without CAD ¹⁰ | No significant association between chronic CAD and severity of dental disease | Findings at variance with those reported previously, ⁹ possibly because of older age of patients in this study |
| 320 veterans ≥ 60 years of age ¹¹ | Oral health parameters more strongly associated with CAD than serum cholesterol levels, body mass index, diabetes and smoking; also, subjects with 1–14 teeth more likely to have CAD than those with 15 or more teeth | No controls included in study |
| 60 patients with acute MI and 60 with chronic CAD ¹² | Periodontal disease may have been associated with acute MI | No controls included in study |
| 166 patients with acute cerebrovascular ischemia and 166 age- and sex-matched neurological controls without stroke who completed a standardized questionnaire; dental examination performed on 66 stroke patients and 60 controls ¹³ | Stroke victims tended to have more severe periodontitis than controls | Unconventional and non-validated assessments of periodontal status; cerebrovascular and cardiac ischemia may be discrete disease processes |
| 151 subjects with either CAD or significant CAD risk factors and 943 healthy controls, 26–53 years of age ¹⁴ | CPITN scores of 4 subjects significantly related to hypercholesterolemia and possibly associated with CAD | Hypercholesterolemia was claimed to link periodontal infections with atherosclerosis, but the cross-sectional nature of this study precludes such interpretations of the data |

MI = myocardial infarction, CPITN = Community Periodontal Index of Treatment Need.

Table 2 Summaries of cross-sectional studies on the relationships between periodontal infections and coronary artery disease (CAD)

| Patient numbers and characteristics | Results | Comments |
|---|--|--|
| Insurance company records of 1,384 men, 45–64 years old ¹⁵ | Weak statistical association between missing teeth and history of ischemic heart disease | Uncertain reliability of missing teeth as surrogate for “sustained oral infections” |
| 88 men referred for diagnostic coronary angiography ¹⁶ | Statistical association between severity of coronary artery stenosis and severity of radiologically evident damage from dental infections | Suggested relationships between oral infections and coronary artery stenosis compromised by unproven reliability of “total dental index” and “pantomographic index” in assessing oral health |
| 5,564 subjects > 40 years old derived from NHANES III ¹⁷ | Significant association between clinically evident attachment loss and self-reported history of MI | Cross-sectional design hampers evidence for a relationship between attachment loss and risk of CAD |
| 12,949 subjects ≥ 18 years old who underwent periodontal examinations and 1,817 who were edentulous (from NHANES III) ¹⁸ | Edentulism and periodontal disease associated with increased systemic inflammatory response, even with adjustment for established risk factors | Cross-sectional design hampers determination of causal relationships |

NHANES = National Health and Nutrition Examination Survey, MI = myocardial infarction.

those already proven for lifestyle changes, in terms of effects on CAD.²⁸ These potential benefits were tested in 4,027 people enrolled in the First National Health and Nutrition Examination Survey (NHANES I), where the prospective cohort trial comprised 2,170 people who were edentulous, with the remainder having periodontitis.²⁶ Over a mean follow-up period of 17 years, 1,238 CAD events (deaths or CAD-related hospital admissions) occurred among the participants. However, after adjustment for the principal risk factors (e.g., smoking, blood pressure and cholesterol level), the risks of CAD were no lower among patients without dental infections (those who were edentulous) than among those with periodontitis. These data argue against a causal relationship between these 2 variables and suggest that the limited associations reported by others may be attributable to other confounding factors, such as smoking. Inadequate prior dental histories (e.g., the prevalence and severity of previous periodontally induced bacteremia) for the patients in this and other studies further compromise more rigorous interpretation of the data. Certainly dental clearances will restore the integrity of epithelial barriers and thereby reduce the potential for recurrent oral sepsis, although CAD may be the chronic problem and may have been initiated while the patient was still dentate.

Can Treatment of Periodontal Infections Reduce the Risk of CAD?

The results of the prospective cohort study on participants in NHANES I did not confirm that treatment for periodontal infections prevents or reverses the prevalence of CAD; in other words, they failed to show that CAD risks were reduced in those with no periodontal infection (edentulous) or exacerbated in those with periodontitis.²⁶ These data therefore suggest that the treatment of dental (especially periodontal) infections should be based principally on dental determinants rather than CAD prevention or treatment. Alternatively, even

if periodontal infections are not independent risk factors for CAD, these 2 diseases might still be associated after due consideration of confounding factors (e.g., smoking). Patients at high risk for CAD may have similar risks for periodontitis, since the diseases have some commonalities in their underlying pathophysiological processes.²⁹ Furthermore, patients may be genetically predisposed to both diseases, in addition to others (e.g., diabetes mellitus). If these relationships are subsequently substantiated by interventional trials, then periodontitis should not be expected to be improved by treatment for CAD, nor CAD arrested or improved after periodontal treatment. Such commonalities may, however, explain the associations between periodontitis and atherosclerosis, diabetes and adverse outcomes for pregnancy identified in animal and population-based studies.³⁰

Conclusions

Current evidence is insufficient to unequivocally support the premise that dental (especially periodontal) infections constitute an independent risk factor for CAD. However, the deposition of atheromatous plaque in the coronary arteries may be associated with many other infections in addition to periodontitis (e.g., bacterial, viral and even parasitic).²⁹ The association between periodontitis and chronic disorders such as diabetes and CAD may also be mediated by either nonspecific (e.g., hypercoagulability, increase in C-reactive protein) or specific (e.g., induction of pathogenic anti-heat-shock protein) mechanisms, in addition to being linked to immunologic components (e.g., serum amyloid protein, fibrinogen). Therefore, evidence for potential associations between dental infections and systemic diseases must be carefully re-examined to distinguish potential confounding factors from other risk factors (e.g., smoking) before treatment with the aim of preventing or treating CAD can be justified. Alternatively, significant similarities in the etiologic and pathogenetic

Table 3 Summaries of retrospective and prospective studies on the relationships between periodontal infections and coronary artery disease (CAD)

| Patient numbers and characteristics | Results | Comments |
|---|--|--|
| Retrospective study | | |
| 10,368 individuals without self-reported CAD and 11,251 with cerebrovascular disease from the Nutrition Canada Survey of 1970–1972 (studied through 1993) ¹⁹ | Significant correlation between periodontal disease and risk of fatal cardiovascular disease | Cross-sectional design hampers determination of a relationship between periodontal disease and risk for CAD |
| Prospective studies | | |
| 214 patients studied for a median of 7.2 years ²⁰ | Statistical evidence for association between oral infections and development of adverse cardiovascular outcomes in patients at increased risk for CAD | Suggested relationships between oral infections and risk of CAD compromised by unproven reliability of “total dental index” and “pantomographic index” to assess oral health |
| 1,147 men 21–80 years old at baseline, derived from Normative Aging and Dental Longitudinal Studies of Boston, followed for 18 years ²¹ | Results support hypothesis that periodontal disease is involved in development of CAD | |
| 22,037 male physicians 40–84 years old in the US Physicians’ Health Study, followed for 12.3 years ²² | Periodontal disease was not an independent predictor of subsequent CAD | |
| 1,372 native Americans with known high risk for diabetes mellitus, evaluated over 10 years ²³ | Alveolar bone level predictive of cardiovascular disease for persons ≤ 60 years old | |
| 22,071 US male physicians followed for a mean of 12.3 years (in randomized, double-blind placebo-controlled trial of ASA and beta-carotene) ²⁴ | Self-reported periodontal disease not an independent predictor of cardiovascular disease in middle-aged to elderly men when adjusted for major cardiovascular risk factors | Lack of precision in self-reported assessments of periodontal disease |
| Prospective cohort studies | | |
| 9,769 adults from the NHANES I, followed for a median of 14 years (Russell’s periodontal index used for assessment of periodontal health) ²⁵ | Periodontal disease and tooth loss weakly associated with CAD development; stronger association for subjects ≥ 50 years old | Russell’s periodontal index scores subject to considerable error |
| 8,032 people from NHANES I Epidemiologic Follow-up Study, followed for a mean of 17 years ²⁶ | Elimination of chronic dental infections did not lead to reductions in risk of CAD | No controlled evidence that dental treatment leads to lower incidence of CAD than among those not treated |
| Prospective questionnaire study | | |
| 44,119 men enrolled in Health Professionals Follow-up Study, contacted biennially by mail over 6 years for self-reported dental history ²⁷ | No overall association between self-reported periodontal disease and CAD Weak association between tooth loss and CAD in those with history of periodontal disease | Self-assessment questionnaires for periodontal disease are imprecise |

ASA = acetylsalicylic acid, NHANES = National Health and Nutrition Examination Study.

processes of these chronic diseases underscore the urgent need for rigorous interventional trials. ➔

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La définition de la santé buccodentaire de l'ADC :

La santé buccodentaire est un état des tissus et des structures associés à l'appareil buccodentaire d'une personne qui contribue à son bien-être physique, mental et social et qui améliore sa qualité de vie, en lui permettant de s'exprimer, de s'alimenter et de socialiser sans douleur, malaise ou gêne.

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